An Engineering Understanding of the Small Intestine

by

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ABSTRACT

The main objective of this research was to understand phenomena occurring during food digestion and nutrients absorption in the small intestine from an engineering perspective. Intestinal flow and mixing processes were simulated using a dynamic *in vitro* Small Intestine Model (SIM). Of particular interest was to study the effect that mixing and food formulation has on glucose absorption and starch hydrolysis.

Results showed the effect of segmentation motion on nutrient delivery to the intestinal wall as a consequence of changes in the mass transfer coefficient. This is most likely due to the increased mixing in the SIM. Experiments of starch digestion with and without the presence of guar gum have shown that viscous fibres reduce the rate of starch digestion and glucose absorption by impairing mixing and reducing diffusion within the fluid. Similarly, use of particulate systems demonstrated a significant effect on the delivery rates. Flow visualization techniques used for studying flow paths in the SIM showed that this *in vitro* model reproduces the characteristic flow events and mixing found in the small intestine *in vivo*.

This research provides insights into the role of mixing on enhancing mass transfer on the course of digestion-absorption processes and also the action of viscous polysaccharides on the delay of glucose absorption in the small intestine. The end findings resulted in a better understanding of the factors which control the development of new functional food that could be applied both in academia and industry.
This project was partially supported by the Programme Alßan, the European Union Programme of High Level Scholarships for Latin America, scholarship number E07D402060MX and by the Mexican National Council for Science and Technology (CONACYT), scholarship number 230576. This PhD was also possible thanks to the support of the National Polytechnic Institute Mexico (CICATA-IPN).
Specially dedicated to my family
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For the Positron Emission Particle Tracking (PEPT) facility, I would like to thank Dr. David Parker, Dr. Thomas Leadbeater and Dr. Joseph Gargiuli. With regards to the use of the Flow visualization laboratory, I would like to thank Dr. Mark Simmons for sharing his expertise and assisting me with the use of the Planar Laser Induced Fluorescence (PLIF) technique I would also like to show appreciation to Ms. Lynn Draper and Ms. Elizabeth Kent for their assistance during my PhD.
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## Nomenclature

- **\( C(x,t) \)**: Drug concentration at any distance along the intestine at time \( t \), mass/m³
- **\( c_1 \)**: Concentration in the bulk solution, mol/m³
- **\( c_{1i} \)**: Concentration at the interface, mol/m³
- **\( C_{\text{inf}} \)**: Concentration at the infinite, mol/m³
- **\( D \)**: Diameter of the pipe, m
- **\( D_{AB} \)**: Diffusion coefficient or diffusivity, m²/s
- **\( D_{\text{mem}} \)**: Diffusion coefficient of nutrient through the membrane
- **\( F \)**: Frequency of oscillation, s⁻¹
- **\( x_o \)**: Amplitude centre-to-peak, m
- **\( J \)**: Diffusion flux of material per unit area, mol/m²s
- **\( K \)**: Overall mass transfer coefficient, m/s
- **\( k_a \)**: Apparent first-order absorption rate constant
- **\( k_{bp} \)**: Local mass transfer coefficient, m/s
- **\( K_m \)**: Michaelis half saturation constant
- **\( k_{rep} \)**: Recipient side mass transfer coefficient, m/s
- **\( L \)**: Characteristic length, m
- **\( l_{\text{mem}} \)**: Thickness of the membrane, m
- **\( N_i \)**: Molar flux at the interface, mmol/m²s
- **\( Nu \)**: Nusselt number
- **\( Re \)**: Reynolds number
- **\( Re_n \)**: Net Reynolds number
- **\( Re_o \)**: Oscillatory Reynolds number
<table>
<thead>
<tr>
<th>Variable</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>$R_{sys}$</td>
<td>Resistance from the system</td>
</tr>
<tr>
<td>$S$</td>
<td>Substrate concentration, mol/m$^3$</td>
</tr>
<tr>
<td>$Sh$</td>
<td>Sherwood number</td>
</tr>
<tr>
<td>$Sh_n$</td>
<td>Sherwood number from net flow</td>
</tr>
<tr>
<td>$Sh_o$</td>
<td>Sherwood number from oscillatory flow</td>
</tr>
<tr>
<td>$S_o$</td>
<td>Initial substrate concentration, mol/m$^3$</td>
</tr>
<tr>
<td>$T$</td>
<td>Time, s</td>
</tr>
<tr>
<td>$v_{ave}$</td>
<td>Average velocity due to segmentation and peristalsis, m/s</td>
</tr>
<tr>
<td>$V_{max}$</td>
<td>Maximal rate of changing</td>
</tr>
<tr>
<td>$v_{osc}$</td>
<td>Oscillatory velocity induced by segmentation, m/s</td>
</tr>
<tr>
<td>$W$</td>
<td>Lambert W function</td>
</tr>
<tr>
<td>$x, y, z$</td>
<td>Position, m</td>
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</table>

**Greek letters**

<table>
<thead>
<tr>
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<th>Description</th>
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<tbody>
<tr>
<td>$A$</td>
<td>Longitudinal spreading coefficient, m$^2$/s</td>
</tr>
<tr>
<td>$B$</td>
<td>Linear flow velocity, m/s</td>
</tr>
<tr>
<td>$\dot{\gamma}$</td>
<td>Shear rate, s$^{-1}$</td>
</tr>
<tr>
<td>$\rho$</td>
<td>Density of the fluid, Kg/m$^3$</td>
</tr>
<tr>
<td>$\nu$</td>
<td>Mean velocity, m/s</td>
</tr>
<tr>
<td>$\mu$</td>
<td>Viscosity of the fluid, Pa.s</td>
</tr>
<tr>
<td>$\mu_{app}$</td>
<td>Apparent viscosity, Pa.s</td>
</tr>
</tbody>
</table>
# Abbreviations

<table>
<thead>
<tr>
<th>Abbreviation</th>
<th>Description</th>
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<tbody>
<tr>
<td>ALP</td>
<td>Alkaline Phosphatase</td>
</tr>
<tr>
<td>ANOVA</td>
<td>Analysis of Variance</td>
</tr>
<tr>
<td>BMI</td>
<td>Body Mass Index</td>
</tr>
<tr>
<td>CCD</td>
<td>Charged Coupled Device</td>
</tr>
<tr>
<td>CFD</td>
<td>Computational Fluid Dynamics</td>
</tr>
<tr>
<td>CMC</td>
<td>Carboxymethyl cellulose</td>
</tr>
<tr>
<td>CT</td>
<td>Computer Tomography</td>
</tr>
<tr>
<td>DNS</td>
<td>3,5-dinitrosalicylic acid</td>
</tr>
<tr>
<td>DPPC</td>
<td>Dipalmitoylphosphatidylcholine</td>
</tr>
<tr>
<td>EE</td>
<td>Entrapment encapsulation</td>
</tr>
<tr>
<td>EPI</td>
<td>Echoplanar Magnetic Resonance Imaging</td>
</tr>
<tr>
<td>ESEM</td>
<td>Environmental Scanning Electron Microscopy</td>
</tr>
<tr>
<td>GI</td>
<td>Gastrointestinal</td>
</tr>
<tr>
<td>IFR</td>
<td>Institute of Food Research</td>
</tr>
<tr>
<td>LUV</td>
<td>Large Unilamellar Vesicles</td>
</tr>
<tr>
<td>MLV</td>
<td>Multilamellar vesicles</td>
</tr>
<tr>
<td>MMC</td>
<td>Migration Motor Complex</td>
</tr>
<tr>
<td>MRI</td>
<td>Magnetic Resonance Imaging</td>
</tr>
<tr>
<td>MS</td>
<td>Microspheres</td>
</tr>
<tr>
<td>MTC</td>
<td>Mass Transfer Coefficients</td>
</tr>
<tr>
<td>MWCO</td>
<td>Molecular Weight Cut Off</td>
</tr>
<tr>
<td>NHS</td>
<td>National Health Service</td>
</tr>
<tr>
<td>Abbreviation</td>
<td>Description</td>
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<tr>
<td>--------------</td>
<td>-------------</td>
</tr>
<tr>
<td>OMTC</td>
<td>Overall Mass Transfer Coefficients</td>
</tr>
<tr>
<td>PAMPA</td>
<td>Parallel Artificial Membrane Permeation Assay</td>
</tr>
<tr>
<td>PC</td>
<td>Phosphatidylcholine</td>
</tr>
<tr>
<td>PE</td>
<td>Phosphatidyl ethanolamine</td>
</tr>
<tr>
<td>PEPT</td>
<td>Positron Emission Particle Tracking</td>
</tr>
<tr>
<td>PET</td>
<td>Positron Emission Tomography</td>
</tr>
<tr>
<td>PI</td>
<td>Phosphatidylinositol</td>
</tr>
<tr>
<td>PIV</td>
<td>Particle Image Velocimetry</td>
</tr>
<tr>
<td>PLIF</td>
<td>Planar Laser Induced Fluorescence</td>
</tr>
<tr>
<td>pNPP</td>
<td>p-nitrophenyl phosphate</td>
</tr>
<tr>
<td>RC</td>
<td>Regenerated cellulose</td>
</tr>
<tr>
<td>RDS</td>
<td>Rapidly Digestible Starch</td>
</tr>
<tr>
<td>RS</td>
<td>Resistant Starch</td>
</tr>
<tr>
<td>SDS</td>
<td>Slowly Digestible Starch</td>
</tr>
<tr>
<td>SEM</td>
<td>Scanning Electron Microscopy</td>
</tr>
<tr>
<td>SGF</td>
<td>Simulated Gastric Fluid</td>
</tr>
<tr>
<td>SHIME</td>
<td>Simulator of the Human Intestinal Microbial Ecosystem</td>
</tr>
<tr>
<td>SIF</td>
<td>Simulated Intestinal Fluid</td>
</tr>
<tr>
<td>SIM</td>
<td>Small Intestine Model</td>
</tr>
<tr>
<td>STR</td>
<td>Stirred Tank Reactor</td>
</tr>
<tr>
<td>TIM</td>
<td>Gastrointestinal Tract Model</td>
</tr>
<tr>
<td>VMD</td>
<td>Volume Mean Diameter</td>
</tr>
<tr>
<td>WHO</td>
<td>World Health Organization</td>
</tr>
</tbody>
</table>
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CHAPTER 1 - INTRODUCTION

1.1 Obesity overview - challenges
According to the latest report from the World Health Organization (WHO) rates of obesity have more than doubled during the past three decades in virtually every country around the world. With approximately 1.5 billion people being overweight and over 375 million adults identified as clinically obese, the problem is reaching epidemic levels with substantial economical implications (WHO, 2011). Consistent with this global trend, the prevalence of obesity has been also found to be dramatically increasing in the UK (Yurgin et al., 2008). Whilst less than 10% of the UK population was classified as obese in 1980, this figure is expected to have grown to 60% by 2050, with an estimated cost to the National Health Service (NHS) of about 6.5 billion pounds per annum (Foresight, 2007). One of the biggest concerns is that this condition is becoming more common in children under the age of five (WHO, 2011). Although several policies are being proposed to reduce and reverse these trends, at present, no country in the world has a comprehensive, long-term strategy to deal with the challenges posed by obesity (Foresight, 2007).

Health implications. Being overweight or obese increases the risk of a wide range of chronic diseases raising levels of morbidity and mortality and reducing the lifespan by up to nine years (Johnstone, 2010). People who are overweight (Body Mass Index, BMI ≤ 25) or obese (BMI > 30) are susceptible to develop depression, type 2 diabetes, hypertension, cardiovascular diseases (stroke) and in some cases even cancer (WHO, 2011, Foresight, 2007). Latest studies also suggest a correlation between obesity and the risk of developing neurological diseases (Scott, 2011, Profenno et al., 2010). Thus, obese people are more susceptible to develop Alzheimer’s disease than slim people (Scott, 2011). Although obesity
can be prevented, the combination of the modern lifestyle, physical inactivity and the availability of high caloric foods in the market makes it difficult to reduce and reverse the incidence of people under this condition.

**Foods in a hectic life.** Nowadays, as a result of lifestyles with less time to prepare or eat our own food at home, most people are eating more highly-processed and easy-to-cook foods; adding saturated lipids and rapidly digested carbohydrates on their daily diets. According to different studies, when rapidly digested carbohydrates and refined sugars are included in diets on a daily basis, blood glucose and insulin responses could be enhanced, resulting in detrimental health effects (Norton et al., 2006, Englyst et al., 1999, Read and Welch, 1985). To avoid this scenario, food and pharmaceutical industries face a new challenge to develop novel food and drug formulations with structural and functional properties to prevent and reduce obesity worldwide.

In an effort to do this, foods with reduced fat and sugars have been designed. These ingredients can be substituted with fibres, modified starches and natural or artificial sweeteners to reduce the caloric content of the food. By using fat replacers, for example, the energy density of foods can be potentially reduced. It has been reported that a reduction in the proportion of fat in the diet by 10% can result in a corresponding decrease of 238 kcal per day of the total energy intake and can produce a weight loss of approximately 3.2 kg (Jones and Jonnalagadda, 2005). Fat replacers on the market encompasses carbohydrates-, protein- and fat-based fat replacers (Jones and Jonnalagadda, 2005).

On the other hand, sugars have been substituted with natural and artificial sweeteners. Natural sweeteners include polyols (sugar alcohols), e.g. sorbitol, xylitol, mannitol or extracts
Chapter 1- Introduction

from plants such as *Stevia rebaudiana* and glycyrrhizin from licorice root. Artificial sweeteners are 100 to 25,000 times sweeter than sucrose with low caloric values, such as aspartame, acesulfame K, sucralosa, neotame and saccharin (Samra and Anderson, 2005). As fat substitutes, the replacement of sugars with artificial sweeteners has a substantial impact on the intake of calories (Norton et al., 2006). For example, the caloric values of the most frequently used sugar replacers range from 1.6 to 3.0 calories per gram whereas sugar and most other carbohydrates provide up to 4 calories per gram, having little or no effect on blood glucose and insulin levels. As a result, a variety of foods with low- or free- fat and sugar can be found in the market.

Furthermore, potassium has been used as an alternative to reduce sodium intake levels in foods to decrease conditions such as hypertension (Gyu-Hee, 2011). Nevertheless, it has a bitter taste that limits its use. Based on that, research has focused on the development of salt substitutes and their applications in foods (Braschi et al., 2009, Horita et al., 2011).

Regardless of all this effort, these products tend to be unsuccessful in the marked because of their flavour and texture (Norton et al., 2006) and their effect on satiety is still controversial (Bellisle and Perez, 1994). In addition, new drugs that can help to regulate appetite control and energy intake might be developed with no side-effects that can limit the efficacy of current treatments.

The above figures emphasize the importance to develop high quality foods with specific health benefits. Nonetheless, these new developments cannot be achieved without a fundamental understanding of the impact that food formulation and structure has on human digestive processes. As a consequence, food digestion and absorption must be evaluated by
considering the physical processes occurring in the gastrointestinal tract. This is a crucial step in the design of food and drug formulations that can deliver nutrients or *active molecules* at specific time and location i.e. controlled and sustainable release.

Through this thesis, the term ‘*active molecules*’ or ‘*actives*’ refers to compounds present in food or drug formulations that have physiological functions; providing health benefits to the consumer. Some examples are vitamins, antioxidant and probiotics. Due to their importance on human physiology, a better fundamental understanding of actives absorption is crucial to develop novel foods for specific applications.

**Structured foods opportunity.** Processed foods could be **structured** to control the rate of release of certain nutrients or slow the rate of stomach emptying, which would limit the amount of food that people could consume (Foresight, 2007). For instance, by adding soluble fibres such as guar gum, β-glucans, pectin and insoluble fibres into food systems it may be possible to reduce the risk of insulin intolerance and type 2 diabetes by changing the absorption rate of molecules such as glucose into the blood (Norton et al., 2006, Brennan, 2005, Blackburn et al., 1984, Jenkins et al., 1978). Minekus et al. (2005) have demonstrated that by adding guar gum to a low fat yogurt, cholesterol levels were reduced. Individual macronutrients could also control the rate of digestive processes influencing satiety and condition the absorption of nutrients (Johnstone, 2010). To do this successfully will however require new approaches to food processing and manufacturing.

Since food formulation has a large influence on the rate of food digestion and absorption, special interest is shown in this project to understand the effect that food formulation has on digestion. This is quite challenging because it is not easy to evaluate the performance of
foods under \textit{in vivo} (in the human body) digestion and absorption processes. These limitations could be overcome by using realistic models \textit{in vitro} that are able to reproduce the physiological conditions found in the gastrointestinal tract, taking into account motility and fluid dynamics of the lumen content. There are similar challenges for the pharmaceutical sector, i.e. the design of oral drug delivery vehicles which effectively carry drugs from the mouth to the small intestine. The interests of food and pharma industries thus coincide.

Overall, the development of new food and pharmaceutical formulations must be driven by a better understanding of the processes by which products are digested and absorbed in the body, combined with knowledge of the release and absorption of the nutrient in the gastrointestinal tract. Since most of the absorption takes place in the small intestine this work is focus on the digestive processes that occur there.

1.2 Mass transfer phenomena in the small intestine

In engineering, molecular and convective transport phenomena can be described by mass transfer. Mass transfer operations have been widely studied in a variety of chemical applications both to separate molecules from one phase to another and to increase the rate and yield of chemical reactions. Mass transfer phenomena are also important in the scale-up of processes and in their control and optimization. Likewise, mass transfer concepts can be also applied to physiological processes. For example, in the intestine actives absorption can be modelled using diffusion and mass transfer coefficients. In this context, the present research combines mass transfer theory with relevant aspects of gastrointestinal physiology to increase understanding of the impact that food formulation and structure has on the digestion and absorption rates inside the body.
1.3 Objectives

The main objective of this work was to understand the physicochemical phenomena occurring during food digestion by using a dynamic \textit{in vitro} Small Intestine Model (SIM). Of particular interest was to study the effect of mixing and food formulation on starch hydrolysis and glucose absorption by:

- Modelling of the processes underway in flow and digestion
- Estimating the mass transfer coefficients between bulk and wall as a function of mixing and food formulation
- Describing flow patterns in the SIM using flow visualization techniques

In addition, an attempt was made to use novel coating structures to reduce the rate of actives digestion and absorption throughout the epithelium.

Release of encapsulated actives was studied under controlled conditions, similar to those found in the small intestine. A development of an efficient way of encapsulating actives was also tested to make the structures.

This work provides insight into the role of mixing on enhancing mass transfer on the course of digestion-absorption processes and also the action of viscous polysaccharides on the delay or reduction of glucose absorption in the small intestine. The end results will be a better understanding of the factors which control the development of new functional food which could be applied both in academia and industry.
1.4 Thesis layout

Chapter 2. Literature review

The chapter presents an overview of the fundamental aspects of the gastrointestinal physiology along digestion and absorption processes. A comprehensive review of the models developed to study these phenomena are also included. As digestion and absorption would be studied from an engineering perspective, relevant concepts of fluid dynamic and mass transfer are briefly discussed in this chapter. The last part is related to the impact that structured foods such as viscous liquids and encapsulated actives may have on the delivery of nutrients in the small intestine.

Chapter 3. Materials and methods

A list of the materials and a description of the common experimental techniques used through this project is included in this chapter.

Chapter 4. Absorption of actives in the SIM

This chapter is related to absorption processes in the SIM, studying mass transfer in single phase liquids. The chapter is divided into two sections related to: (i) the impact of mixing on mass transfer coefficient and (ii) the effect that wall movements or segmentation contractions (frequency) has on glucose absorption. To visualize the effect of food formulation and mixing on glucose absorption techniques such as Planar Laser Induced Fluorescence (PLIF) and Positron Emission Particle Tracking (PEPT) were also used (Chapter 6).

Chapter 5. Digestion and absorption in the SIM

This chapter describes a set of experiments that include both chemical digestion and actives absorption in the SIM. The main objective of this chapter was to investigate the effect of
Chapter 1- Introduction

increasing viscosity of digesta on starch hydrolysis and glucose absorption. Furthermore, parameters such as enzyme and substrate concentration were studied to evaluate starch kinetics within the model system.

Chapter 6. Mixing processes in the SIM

A description of the experiments performed with PLIF and PEPT to study the fluid flow inside the SIM is included in this chapter. Parameters such as viscosity and mixing were investigated. PEPT has also been used to determine the velocity field in the system and suggest what velocities of lumen content may occur (using a free tracer particle).

Chapter 7. Encapsulation of actives

The seventh chapter covers the production of micro- and nano-capsules for controlled release of encapsulated actives. The first section focuses on gelled particles, including morphology, entrapment efficiency and in vitro release whereas the second section describes the results of producing multilamellar liposomes for control release of proteins. The physicochemical properties of the standard liposomes are compared with gelosomes for stability and control release.

Chapter 8. Conclusions and future work

The conclusions and recommendations for future work are summarized in Chapter 8.

1.5 Publications and Conferences

The results from this work have been published and presented at different forums as described below.


Publications


Conferences


CHAPTER 2 - LITERATURE REVIEW

2.1. Introduction

As mentioned in Chapter 1, predicting the extent of food digestion and absorption in the gastrointestinal tract is essential to the selection and development of structured foods that are inherently healthier. To predict the *in vivo* performance of food after ingestion, it is crucial to establish the physiological and physicochemical factors affecting chemical reaction and absorption processes in the human body. This chapter describes the literature related to digestion and absorption in the gastrointestinal (GI) tract and the physicochemical phenomena involved from an engineering perspective. Firstly, an overview of the GI tract is presented. Special interest is given to the physiology of the small intestine, describing the fluid dynamics and mixing processes taking place in it. Food digestion and actives absorption principles are also included, followed by a description of the models developed to study these phenomena *in vivo, in vitro* and *in silico*. Secondly, as the small intestine can be considered as a tubular bioreactor in which enzymatic reactions occur, an overview of the basic principles of fluid flow and chemical reactions is given. Finally, a description of how food structure and formulation influence the rate of digestion and absorption in our body is included; highlighting the effect that encapsulation techniques and encapsulated structures have on the release of active components.

2.2. Gastrointestinal Tract overview

Details of the physiology of the GI tract can be found in Guyton and Hall (2006), Levy et al. (2006) and Widmaier et al. (2006), and from which most of the discussion below has been taken. Few studies have considered the role of the GI tract in an engineering way, and these will be discussed later. The human GI tract is a highly regulated multistage system in which
solid/liquid foods are processed, dissolved and broken down into small molecules that can be easily absorbed through the epithelial cells and then into the bloodstream. Once ingested, food undergoes a series of unit operations that allow the bodies’ energy requirements to be met. These operations take place in four distinctive reactive sections of the GI tract; mouth, stomach, small and large intestine, as shown in Figure 2.1.

Each compartment has a specific function and similar purpose; to maximize digestion and absorption in the GI tract. The next section presents a brief description of the GI tract organs and processes involved in food digestion.

**Figure 2.1** Scheme of the gastrointestinal tract.

### 2.2.1 Mouth

Food digestion starts in the mouth (Figure 2.2), an oral cavity, in which foods are reduced in size by the teeth through chewing cycles that increase the surface area available for digestion. The number of chewing cycles not only varies among individuals but also according to the food characteristics and structure (van der Bilt, 2009). During mastication, food particles are
mixed with saliva produced by three pairs of major glands: parotid, submandibular and sublingual glands (Figure 2.2).

**Figure 2.2** Diagram of the side view of the structure of the mouth.

Saliva, a complex dynamic biological fluid, is mainly composed of water (99.5%), proteins (0.3%) and inorganic and trace substances (0.2%) (van Aken et al., 2007). Proteins and inorganic components in saliva are mainly: glycoproteins such as mucins, proline-rich glycoproteins; enzymes (α-amylase, carbonic anhydrase); immunoglobulins; peptides (cystatins, statherin, histains) and electrolytes (sodium, potassium, chloride and bicarbonate). Secretion of saliva ranges from 0.3 to 7.0 ml of saliva per minute with about 0.5 - 1.5 litres of saliva secreted per day and a pH from 6.2 to 7.4 (van Aken et al., 2007). Saliva is a non-Newtonian fluid and its viscosity decreases upon increasing shear rate. Rheological properties of saliva are consistent with the behaviour of a weak gel in which a liquid, gaseous and gel phases coexists simultaneously (van Aken et al., 2007). This behaviour stimulates flavour perception and lipid melting phase occurring at the first stages of mastication (Norton et al., 2006, Taylor and Linforth, 1996).

During mastication, water present in saliva moistens and dissolves the food to help further digestion whereas mucus composed of mucins functions as a glue to bind food tiny particles
into a smooth paste called ‘bolus’ that can be easily swallowed. Salivary amylase, on the other hand, starts the digestion of the starch contained in food.

Once swallowed, the bolus is transported into the stomach through the pharynx and oesophagus by peristaltic waves. Although these elements of the GI tract do not contribute directly to the digestion process, they play an important function in transporting the bolus from the mouth to the stomach.

### 2.2.2 Stomach

The stomach functions as a storage vessel in which the bolus is partially ground and combined with digestive juices secreted by gastric glands. Anatomically, the stomach can be divided into four major sections: fundus, body, antrum and pylorus (Figure 2.3).

![Diagram of the stomach showing the different sections](Widmaier et al., 2006)

**Figure 2.3** Diagram of the stomach showing the different sections (Widmaier et al., 2006).

The fundus and the body integrate the proximal part of the stomach, where large volumes of food (1.5 - 4.0 litres) can be stored for long periods of time (3 - 4 hours) (Kong and Singh, 2008a). As long as food is in the stomach, peristaltic constrictor waves occur in the proximal region of the stomach. Nevertheless, these waves are normally weak, occurring once every 15
or 20 seconds, i.e. much of the gastric contents remain unmixed for long periods as shown by the echoplanar magnetic resonance images (Marciani et al., 2001). The antrum, or distal stomach, functions as a grinder-mixer-pump. Here contractions are vigorous, compressing the food several times into the stomach walls, reducing food particle size and mixing the contents of the stomach by ‘retropulsion’. The resulting mixture of food and gastric juices is called “chyme”; a semi fluid paste with varied rheological properties that contains molecular fragments of proteins, polysaccharides and fat droplets (Widmaier et al., 2006). Finally, the antral section of the stomach propels chyme into the upper part of the intestine by means of the pylorus through the pyloric valve (Figure 2.3).

Motility in the stomach is not only fundamental for mixing enzymes and substrates but also to control the passage of chyme. The residence time of food inside the stomach depends on its composition and physical properties (Kong and Singh, 2008a). In the stomach foods are ‘classified’ according to their density or particle size: liquids are located in the antrum whereas solid particles and lipids are mainly placed in the body of the stomach (Schulze, 2006). That means that the stomach serves like a sieve where liquid and small particles leave the stomach more quickly than larger ones. Large particles move through the stomach more slowly than smaller ones, since they have to be small enough, less than 2.0 mm, to pass the pyloric valve between the stomach and the next compartment of the GI tract: the small intestine (Hunt, 1983). In healthy individuals, the half-emptying time for non-nutrients liquids is around 20 minutes (Camilleri, 2006). In contrast, solids are retained selectively in the stomach for “churning” or trituration by the high shearing forces set up by contractions in the antrum as solids are propel toward the closed pylorus. As result of trituration, solid foods are emptied after their size has been reduced to 2.0 mm (Camilleri, 2006).
Chapter 2 - Literature review

Gastric emptying into the intestine is regulated in response to the nature of duodenal contents and is influenced by several factors including the volume, consistency and fat content. Gastric emptying of solids is a two-phase process: an initial lag or retention period followed by a generally linear postlag emptying phase (Camilleri, 2006). The stomach empties solids completely after approximately 3 - 4 hours. Motor and digestive processes are modulated by several hormones, including gastrin, cholecystokinin (CCK) and glucose-regulating hormones such as insulin, glucagon and incretins (Camilleri, 2006). Gastric content is delivered to the duodenum through a negative feedback mechanism mediated by duodenal receptors (Kong and Singh, 2008a) at a rate of 200 kcal per hour (Hunt, 1983, Camilleri, 2006). Although controversial, it has been reported that increasing the viscosity of liquid meals delays gastric emptying and increases satiety (Kong and Singh, 2008a).

Within the stomach, protein and lipid digestion starts as a result of pepsins and gastric lipases respectively. Meanwhile, starch digestion by salivary amylase continues until the acidic environment of the stomach inhibits the hydrolytic action of this enzyme - that can be up to an hour after food enters the stomach. Gastric secretion (2 - 3 litres per day) contains about 0.8 to 1.0 mg/ml pepsin and about 1.5 mg/ml of mucin (Kong and Singh, 2008a). Furthermore, about 2.0 L of hydrochloric acid are also secreted in the stomach per day increasing the concentration of hydrogen ions in the lumen of the stomach up to ~150 mM (Widmaier et al., 2006). Secretion of HCl causes a reduction in the pH of the stomach and food proteins unfolding. Gastric secretion is a viscous fluid with viscosity in the range of 0.01 to 2.0 Pa.s and density close to the density of water. Under fasten conditions, the rate of gastric secretions is approximately 1.0 mL/min, increasing from 10 to 50 mL/min immediately after food digestion (Kong and Singh, 2008a). Gastric pH varies from 2.0-
6.5/7.0 depending on food and phase of gastric emptying. Increase in food amount, protein content and meal viscosity increases secretion.

2.2.3 Small intestine

The small intestine, the focus of this thesis, is responsible for most of the digestion and absorption through the epithelial cells that line it’s surface. Comprised of the duodenum, jejunum and ileum (Figure 2.4), the small intestine is the longest part of the GI tract, reaching a length between 2 and 6 meters (Guyton and Hall, 2006). The external diameter of the gut is on average 2.5 cm, varying from 5 cm in the upper portions of the duodenum to less than 2 cm in the distal ileum (Stoll et al., 2000). The duodenum, the upper part of the small intestine, is a tubular organ with muscular walls in which chyme is mixed with biliary and pancreatic secretions from the liver and pancreas (sections 2.2.5 and 2.2.6), employed for digestion and emulsification of lipids, respectively. Normally, most of the chyme is digested and absorbed in the first segments of the small intestine, in the duodenum and jejunum (Widmaier et al., 2006).

Within the duodenum, the acidic chyme (pH ~2.5) from the stomach is initially neutralized with sodium bicarbonate from the pancreas to avoid denaturalization of the digestive
enzymes. Once neutralized (pH ~ 6.5), the chyme is mixed with hydrolytic enzymes present in the pancreatic secretions: proteases, amylases and lipases that continue the digestion of proteins, carbohydrates and lipids, respectively. In the small intestine, approximately 8.0 L of ingested and secreted water enter per day, however, only 1.5 L pass on the large intestine because almost 80% of the fluid is absorbed in the small intestine (Widmaier et al., 2006). All these processes are highly regulated by hormones and neural mechanisms (Levy et al., 2006, Widmaier et al., 2006).

As in the stomach, motility enhances chemical reactions and brings partially digested molecules such peptides and oligomers to the brush border of the intestinal wall where digestion is completed by the enzymes present on the intestinal wall. These biopolymers are broken down into simple molecules; aminoacids and monosaccharides for further absorption. Contractions of the duodenum typically follow gastric antrum contractions, preventing regurgitation of duodenal contents into the stomach. The flow and mixing of chyme in the intestine, the main area of interest of this research, is determined by the movements of the muscle layers described in detail in section 2.3.1. Analysis of this flow should be done by formulation of the equations that govern muscle movement and chyme rheology (Macagno and Christensen, 1980).

The small intestine transport solids and liquid at approximately the same rate. As a result of the lag phase for the transport of solids from the stomach, liquids typically arrive in the intestine before solids. However, it takes about 150 minutes for half the solid and liquid chyme of similar caloric density to traverse the small bowel (Camilleri, 2006). It is believed that complex carbohydrates or fat that reach the distal part of the small intestine serves as feedback control of proximal intestinal motility (Camilleri, 2006).
2.2.4. Large intestine

The last stage of digestion is the large intestine (colon) in which the remaining water from indigestible food matter is absorbed and unusable food matter (wastes) is stored and finally eliminated from the body.

Figure 2.5 shows the anatomical structure of the large intestine. The major sections of the large intestine consist of ascending, transverse, descending, and sigmoid portions. The ileocelecal valve of the ileum in the small intestine moves material into the large intestine at the caecum. The portion of the ileum containing the ileocelecal valve projects slightly into the caecum. Increases in colonic pressure squeeze the valve shut, whereas increases in ileal pressure open it, therefore effectively preventing reflux of colonic contents into the ileum (Guyton and Hall, 2006). Material passes through the ascending, transverse, descending and sigmoid portions of the colon and finally enters into the rectum where the undigested material is excreted from the body. The large intestine removes about 90% of the fluid, it converts 1000-2000 ml of isotonic chyme that enters it each day from the ileum, to about 200 - 250 ml of semisolid waste.

Figure 2.5 Diagram of the large intestine
Accessory organs of the GI tract

Digestive secretions such as saliva, salts and the enzymes required by the GI tract are produced by accessory organs: salivary glands, liver, gallbladder and pancreas as discussed below.

2.2.5 Liver and gallbladder

The liver (Figure 2.6), located in the upper right portion of the abdomen is in charge of a variety of endocrine and exocrine functions. As only the latter are directly involved in gastrointestinal functions, this section focuses on the exocrine functions of the liver. The hepatic function most important to the digestion is the synthesis and secretion of bile. Bile contains bicarbonate ions, cholesterol, phospholipids, bile pigments, and most important for digestion, bile salts; essential for both digestion and absorption of lipids. Between meals, bile is stored in the gallbladder, a small sac underneath the liver that branches from the common hepatic duct (Widmaier et al., 2006). The gallbladder is an active storage vessel of approximately 8 cm lengthwise and 4 cm in diameter (Figure 2.6), which absorbs mineral salts and water received from the liver and converts it into a thick, mucus substance called "bile", to be released when food is present in the stomach (Levy et al., 2006). During digestion, the smooth muscles in the gallbladder wall contract, causing the concentrated bile solution to be injected into the duodenum via the common bile duct, an extension of the common hepatic duct (Figure 2.7) (Widmaier et al., 2006, Guyton and Hall, 2006).
2.2.6 Pancreas

The pancreas is an elongated gland of 12.5 – 15.0 cm of longitude located behind the stomach. Like the liver, pancreas has both endocrine and exocrine functions. The exocrine portion of the pancreas secretes into the duodenum digestive enzymes (Levy et al., 2006) and a fluid rich in bicarbonate ions via the pancreatic duct. From the pancreas amylase is secreted via the pancreatic duct and then common bile ducts (Figure 2.7) into the duodenum where it plays an important role in digestion of complex carbohydrates.
Since the small intestine is a crucial step in the digestion and absorption of essentially all dietary organic molecules, this work is focused on reproducing the physical and chemical phenomena occurring in the human small intestine. The following section describes in detail the physiological digestive and absorptive functions of the small intestine and the fluid dynamics occurring there as a result of propulsion and mixing movements during food digestion.

2.3 Food digestion and active absorption processes

Food digestion involves two types of processes: (i) mechanical digestion, related to the motility process that promote food mixing and displacement thought the GI tract and (ii) chemical digestion, responsible for the catabolic reactions that breakdown proteins, lipids and carbohydrates into small molecules that could be absorbed by the cell membranes.

2.3.1. Mechanical digestion: propulsion and mixing in the small intestine

Fluid motion within the small intestine is generated by three movement patterns: (i) peristalsis, (ii) segmentation and (iii) pendular movements (Macagno and Christensen, 1980). Peristalsis and segmentation are basic electrical rhythm movements that cause contractions of the circular muscle layer of the intestinal wall. Peristaltic contractions spread along down the intestine whereas segmentation contractions are stationary and local. Pendular movements, on the other hand, cause retropulsion of lumen contents with a characteristic back and forth pattern. Peristalsis and segmentation will be reviewed in detail in sections below.

2.3.1.1 Peristalsis

Peristalsis is the progressive contraction of successive sections of circular smooth muscle. Peristaltic waves occur in the small intestine involving a very short segment of intestine.
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Slow waves and action potentials determine the frequency and force of intestinal contractions (Levy et al., 2006). Peristalsis propels chyme through the small intestine in about 3 - 6 hours (Liu et al., 2003) with an average velocity of 1.0 cm per minute (Guyton and Hall, 2006).

2.3.1.2 Segmentation

Mixing contractions or segmentation occur when a section of the small intestine becomes filled with digesta; as a result, a localized concentric contraction takes place in the gut as shown in Figure 2.8 (Guyton and Hall, 2006). This process is repeated at spaced intervals along the intestine and is responsible for the fluid dynamics of its contents. Segmentation, unlike peristalsis, produces mixing of the intestinal content with little net flow. Consequently, segmental contractions create a continuous division and subdivision of the intestinal content modifying concentration gradients of any substance in the chyme (Macagno et al., 1982). Although these type of contractions are only a few cm long and the contraction last a few seconds (Widmaier et al., 2006), the effect that mixing has on mass transfer phenomena occurring in the intestine may be significant. Furthermore, nutrient absorption could be modified in different ways, increasing, for example, as a result of the intestinal wall movements (Nguyen et al., 2008, Udaykumar et al., 2006, Macagno et al., 1982). Intestinal contractions create turbulence and convection currents, which mix the luminal content bringing material from the centre of the lumen to the epithelium (Macagno et al., 1982, Edwards et al., 1988). Actives have to diffuse across a thin relatively unstirred layer of fluid lying adjacent to the epithelium (Blackburn et al., 1984).
In humans, these regular slow waves occur all along the small intestine at a frequency of 11 - 13 per minute in the duodenum which declines along the length of the small bowel to a minimum of 8 - 9 per minute in the terminal part of the ileum (Levy et al., 2006). The slow waves may be accompanied by bursts of action potential spikes. Action potential bursts are localized to short segments of the intestine, promoting the highly localized contractions of the circular smooth muscle that cause segmentation. Major mixing and propulsive movements of the small intestine take place as a consequence of action potentials that stimulate much stronger contractions of the smooth muscle (Levy et al., 2006).

2.3.2 Chemical digestion in the small intestine

As chyme enters the small intestine and flows axially down, macromolecules such as proteins, lipids and carbohydrates are digested (into amino acids, fatty acids and monosaccharides, respectively) by the action of enzymes present in the gastrointestinal secretions as described in following sections.
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2.3.2.1. Protein digestion

Most protein digestion occurs in the stomach and the upper part of the small intestine. In the stomach digestion of proteins is accomplished by the acidic enzyme pepsin within a working pH range from 3.3 to 5.0. Pepsin is secreted into the stomach in the form of an inactive precursor (pepsinogen). Exposure to low pH, by secretion of hydrochloric acid in the lumen of the stomach causes conversion of pepsinogen to active pepsin (Widmaier et al., 2006). Pepsin secretion in adults has been estimated between 20 - 30 kUnits of enzyme activity at 37 °C and 24 hours (Wickham et al., 2006). Although gastric pepsins are essential for the digestion of proteins, this step only provides 10 to 20% of the total protein digestion (Guyton and Hall, 2006). More extensive hydrolysis of proteins occurs in the upper small intestine where protein digestion continues under the action of proteases from the pancreatic secretions. The most important proteases segregated into the small intestine are: (i) trypsin and chymotrypsin which cleave the internal peptidic bonds of proteins, split them into small polypeptides and (ii) carboxypolypeptidase that produces individual amino acids from the carboxyl ends of the polypeptides (Guyton and Hall, 2006). At this stage, the main products of the proteins are di- and tri-peptides that in turn are converted into amino acids by the peptidases present on the brush border of the small intestine (Figure 2.9).

The efficacy of digestive peptidases in the GI tract depends on protein microstructure and its molecular interactions with other food constituents (Lundin et al., 2008). For example, in the case of milk proteins, the open structure of sodium caseinates allows pepsin and trypsin access to target residues, resulting in rapid proteolysis and faster uptake and absorption of amino acids during digestion (Guo et al., 1995). In contrast, the compact globular structure of β-lactoglobulin inhibits enzyme access to potential cleavage sites and therefore the protein is considerably more resistant to hydrolysis. However, changes in conformation and structure
induced by heating or changes in pH make β-lactoglobulin susceptible to peptic hydrolysis (Guo et al., 1995). Precipitation of casein around the isoelectric point (4.5) has been reported to result in delayed gastric emptying, while β-globulin, which is not subject to acid precipitation, empties rapidly into the duodenum (Lundin et al., 2008).

Moreover, molecular interactions between proteins and carbohydrates induced by enzymes and Maillard reactions may also affect the rate at which proteins are digested in the stomach and intestine (Lundin et al., 2008). Overall, the rate of protein digestion in the GI tract is influenced by several factors that must be taken into account to design structured foods that could impact on satiety and weight regulation.

2.3.2.2 Lipids

Digestion of lipids, triglycerides in a normal diet, starts by the action of lingual and acid-stable gastric lipases secreted in the mouth and the stomach, respectively. Once in the duodenum, lipids are converted into 2-monoglycerides and fatty acids by the action of pancreatic lipase in the presence of surface active bile salts (Guyton and Hall, 2006). During food digestion, between 200 and 250 mg of pancreatic lipase are secreted to convert all the ingested fat (Lundin et al., 2008).

Since lipids are slightly soluble in water they have to be ‘pre-processed’ before they can interact with the pancreatic lipase. As such, a crucial step for fat digestion involves the mechanical breakdown of fat globules into small particles that the water-soluble digestive enzymes can act on. This process is called emulsification of the fat, and it begins in the stomach by mixing the fat with the products of stomach digestion. Then, most of the emulsification occurs in the duodenum under the influence of bile from the liver. Bile
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contains a large quantity of bile salts as well as the phospholipid lecithin that form micelles with lipids. Once fatty acids are released and emulsified with bile, they can cross the epithelium for absorption.

As with proteins, the rate of lypolysis is highly influenced by the structure of the emulsion formed in the stomach, one of the crucial factors on fat digestion is the emulsion surface area (Lundin et al., 2008). For example, in gastric aspirates the lypolysis of a fine emulsion was almost three-fold faster than a coarse emulsion (Armand et al., 1999). Understanding the interaction of emulsions with gastrointestinal environments will enable to develop and design emulsions with specific structures and health benefits in vivo.

2.3.2.3 Carbohydrates - main source of energy

Major sources of carbohydrates within a normal human diet are; sucrose, a disaccharide from the cane sugar, lactose from milk and starch from cereals, roots and tubers (Guyton and Hall, 2006). Starch can represent up to 80% of the carbohydrates present in staple foods of population around the world. Since starch is one of the most important glycemic carbohydrate components in foods, this section presents a comprehensive overview of its digestion.

As described in section 2.2.1, digestion of starchy food begins in the mouth when particle food size is reduced and mixed with amylase from the salivary glands. This process continues until food bolus is mixed with the hydrochloric acid secreted into the stomach inactivating the salivary α-amylase and stopping the reaction. When chyme enters to the duodenum, a more powerful α-amylase in the pancreatic secretions continues the hydrolysis of starch in the small intestine. The pancreatic α-amylase is an endoglycosidase with a molecular weight of 54000 Daltons that cleaves the internal α-1,4 links of the starch structure (Dona et al., 2010).
α-Amylase secretions in adults has been reported between 5 to 15 nM with high activities (Slaughter et al., 2001). The main products of the hydrolysis of starch are: maltose, maltotriose and α-limit-dextrins. These oligosaccharides are completely hydrolysed as glucose by enzymes present on the brush border (Figure 2.9) of the intestinal surface mainly: maltase-glucoamylase and sucrose-iso-maltase.

Digestion of starch could be studied as a solid-liquid two phase reaction (Zhang et al., 2009), where amylase must (i) diffuse into the food matrix, (ii) bind the substrate, and (iii) cleave the glycosidic linkages of the starch molecules (Leloup et al., 2004). Therefore, any factor affecting enzyme activity such as quantity, different isoforms, inhibitors, and the susceptibility of substrate to hydrolytic enzymes will influence starch digestion and the resulting nutritional properties of the material. Factors affecting the binding of amylase to starch will also slow down the enzyme reaction. As such, susceptibility of starch hydrolysis to amylase depends on physical features (Lehmann and Robin, 2007, Englyst et al., 1992), chemical modifications (Chung et al., 2006, Wolf et al., 1999) and processing degree (Anguita et al., 2006, Chung et al., 2006). Physicochemical characteristics of starch such as morphology and particle size, are critical factors on starch digestion and as a consequence, on glucose absorption (Li et al., 2004, Snow and O'Dea, 1981). Generally, α-1,6 glycosidic bonds are more difficult to cleave compared with α-1,4 glycosidic, therefore more α-1,6 glycosidic bonds slow starch digestion (Leloup et al., 2004). Amorphous or completely dispersed starches are highly susceptible to enzyme action. On the other hand, starches that are physically entrapped, such as B-type, raw starch granules, retrograde amylose and chemically modified starches are resistant to digestion (Leloup et al., 2004). Starches may have an intermediate slow digestion pattern if either some degree of physical structure is retained during food processing or structural aspects slow kinetics of digestion.
Starch digestion rates in particular have been directly correlated to postprandial glucose and insulin responses (Zhang et al., 2009). Rapidly absorbed sugars and carbohydrates, such as sucrose, result in high glucose blood levels (Read and Welch, 1985). Healthy people are able to cope with this condition by adjusting their insulin levels as required. Nevertheless, people with diabetes either do not produce insulin or are resistant to its action. In the first case, they need to inject it (type 1 or insulin-dependent diabetes mellitus) while in the second, they can control their glucose levels with specific diets (type 2 or non-insulin-dependent diabetes mellitus). In each case they cannot deal effectively with a rapid rise in blood glucose.

During the passage of foods through the gastrointestinal tract, there are other factors that affect food digestion and absorption including gastric emptying, enzyme inhibitors and viscosity in the digestive tract affecting enzymes accessibility to substrate and the absorption of glucose. It is possible that these can be used to generate a slow digestion effect (Zhang et al., 2009). Overall, the rate at which chemical reactions occurs in the gastrointestinal tract determinate the release of nutrients from the complex matrix of the digesta.

2.3.3 Absorption of active molecules

Absorption is related to the transport of nutrients or active molecules through the epithelium (epithelial cells) to the blood stream. Most of the absorption of water and nutrients including vitamins and electrolytes occurs in the small intestine. The absorption surface area of the small intestine is roughly 250 m$^2$, as a result of the corrugations of the intestinal wall to maximize absorption. The intestinal mucosa is a physical and biochemical barrier to absorption, composed by the three distinctive features as shown in Figure 2.9.
Wall structure. The small intestinal wall consists of the serosa, the muscularis externa (longitudinal and circular muscle layers), the submucosa and the intestinal mucosa (Figure 2.9). The latter represents the main barrier of the intestinal wall of nutrient and drug uptake (Deferme et al., 2008). Four layers encompass the intestinal mucosa: (i) surface monolayer, (ii) basal membrane, (iii) lamina propria and (iv) lamina muscularis mucosa. The lamina propria layer is composed by connective tissue, nerves, macrophages, mass cells lymphatic and blood vessels that serves to the transport of substances around the body (Deferme et al., 2008).

The inner surface of the small intestine is organized into circular or mucosal folds which not only increase surface area but are hypothesised to act as baffles, mixing the lumen contents (Peters and Brain, 2009). Extending from the luminal surface into the lumen of the small intestine are fingerlike projections known as villi. The surface of the each villi is covered with a layer of epithelial absorptive cells (enterocytes) whose surface membranes form small projections called microvilli (brush border).
The luminal plasma membrane of the absorptive epithelial cells is a selective barrier to the passages of molecules. Membrane regulates transport of active compounds in and out the membranes surrounding cell organelles (Levy et al., 2006). The transport of molecules across the epithelial cells located around the villus of the small intestinal wall is a complex process involving some of the mechanisms described below.

**Main mechanisms of transport occurring in the plasma membrane**

Transport across the epithelial cells of the small intestine surface can be via different mechanisms, such as the following (Duchateau and Klaffke, 2009):

1. **Passive diffusion**, mechanism driven by a concentration gradient, applicable for small and lipophilic molecules, e.g. fructose.
2. **Carrier-mediated transport**, for small hydrophilic molecules. Since a carrier protein is involved, this system is saturable and dependent on a sodium ion gradient. Examples: luminal monosaccharides such as glucose and galactose are transported by this mechanisms across the entocyte brush border (Camilleri, 2006).
3. **Endocytosis**, uptake of particles or large macromolecules. It also requires energy, is slow and limited in capacity, e.g. larger polypeptides.
4. **Paracellular transport** across the tight junctions between adjacent epithelial cell characteristic of hydrophilic molecules (Peters and Brain, 2009).

Hydrophobic and lipophilic molecules are absorbed through the lipid bilayer and the membrane-bound protein regions of the cell membrane located on the microvilli of the small intestine (Peters and Brain, 2009). The mechanisms of membrane transport described above are illustrated in Figure 2.10.
Overall, to absorption of active molecules occurs, four characteristic steps are followed: (i) release of the active from the matrix, (ii) dissolution or dispersion of the active into the GI fluids intestinal content (iii) possible digestion or degradation and (iv) final absorption or complex binding. Amorphous materials tend to show much higher aqueous solubility than crystalline forms of the same compound and different crystal modifications of the same molecule (Johansoon and Paterson, 2008). The solubility of a compound may also be affected by physiological factors such as pH, lipid, and bile acid concentrations. Absorption rates then depend on food properties and physiological factors. In order to predict absorption all these factors must be taken into account to obtain representative data.

Although digestion and absorption processes are not easy to simulate, in vitro and in silico models have been used to study the effect of food formulation in the GI tract.

2.4 Models for studying digestion and absorption

In an effort to reproduce digestion and absorption processes in the GI tract, in vitro and in silico models have been developed. In this section, the principles of physiological-based in
vitro and in silico models for the study and prediction of the human digestion and absorption phenomena are briefly described. Since a wide variety of models have been developed, the discussion is limited to models that could be relevant to this work. A review of the in vivo studies used to evaluate motility and mixing in the GI tract is also presented.

2.4.1 In vivo studies

To evaluate the efficacy of newly developed delivery systems, the most accurate results are obtained indeed from in vivo feeding methods, using humans and animals (Hur et al., 2011). For instance, the disintegration of food in the GI tract can be directly monitored in humans by aspirating fluids from stomach and upper small intestine through a naso-gastric or naso-jejunal tube. Nevertheless, only fluid foods can be tested by this method. Naso-gastric monometric catheters can be also used to monitor gastric motility, however, this study is limited and the intubation procedure is invasive and may cause discomfort to the subjects. As such, non-invasively imaging techniques such as Scintigraphy, Ultrasonography and Echoplanar Magnetic Resonance Imaging (EPI) have been employed to visualize and monitor gastrointestinal functions (Kong and Singh, 2008a).

Scintigraphy is a diagnostic technique used in nuclear medicine, wherein the emitted radiation of radioisotopes or radiopharmaceuticals, are recorded by external gamma cameras in a form of two-dimensional images. Being non-invasive, scintigraphic techniques have been used to evaluate the effects of gastric surgery or autonomic neurophaty (Spiller et al., 2009). These techniques have allowed studying multiphase meals to evaluate gastric emptying of liquids and solids. However, one of the limitations is that subjects could be exposed to substantial doses of ionising radiation.
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*Ultrasonography* is an ultrasound-based diagnostic imaging technique used for visualizing subcutaneous body structures including tendons, muscles, joints, vessels and internal organs. Ultrasonographic techniques are limited to views of the antrum because assessment of the whole gastric volume is difficult due to air/fluid interfaces, which disrupt the ultrasound beam (Marciani et al., 2000).

*Magnetic Resonance Imaging (MRI)* and *Computer Tomography (CT)* scans can produce deep tissue images making these techniques widely used to study gastric motility. Echoplanar Magnetic Resonance Imaging (EPI) is an ultrahigh speed MRI used to study gastric emptying, motility in the gastric lumen and to obtain three-dimensional images of the intragastric distribution of water-containing meals (Marciani et al., 2000). Recent developments in functional EPI techniques have allowed *in vivo* visualisation of a range of formulations (Marciani et al., 2001). For instance, images of alginate beads in the stomach have been presented by different authors (Rayment et al., 2009, Hoad et al., 2009). Hoad et al. (2009) investigated the effect of model solid foods on satiety and reported that gastric emptying was delayed by the presence of gelled beads used as a model.

One of the advantages of EPI is that can be adapted to monitor the viscosity of gastric contents as their dilution by gastric secretions (Schulze, 2006, Kong and Singh, 2008a, Marciani et al., 2001). For instance, when the viscosity of test meals modified with locust bean gum (0.25 - 1.5g/100g) was monitored *in vivo* by the EPI technique; it was found that the viscosity of the meals decreased after ingestion due to dilution with saliva and gastric secretions. For instance it was found that by adding 1.5g/100g meal, the viscosity of the meal fell from 11 to 2 Pa.s. In contrast, the less viscous meal (0.25g/100g) had a much smaller absolute fall (Marciani et al., 2000).
Other imaging techniques used to monitor gastric emptying and motility includes: electrical impedance tomography and manometry. Electrical impedance tomography is an inexpensive technique used in high-conductivity meals to monitor gastric motility (Spiller et al., 2009). On the other hand, the migration motor complex (MMC) of the stomach in the fasting state has been studied by manometry measures pressure; nevertheless with viscous meals the ability to detect movement of the gastric wall is impaired (Spiller et al., 2009). Although these techniques have limitations, they provide useful information on the pattern of flow produced by peristalsis and segmentation movements in the small intestine. According to Imam et al. (2004), when techniques such as videofluoroscopy, manometry, and multiple intraluminal impedance are combined, patterns of flow in the small bowel can be obtained with a good resolution. In addition, fluorescent nanotubes attached to a medication have been used to evaluate the accuracy of the drug in reaching its target whereas the internal organs can be imaged simultaneously (Bergeron, 2011). This is a real time technique that showed promise to study the performance of an active molecule in the GI tract.

Although unanswered questions remain, the last developments in flow visualization imaging techniques will enable a better understanding of gastrointestinal motor functions in future, especially regarding the small intestine in which assessment of flow is difficult.

A significant problem is that in vivo studies and experiments are expensive, time consuming and ethical constraints can limit their use in human experimentation. However, they are critical in establishing accurate parameters that have to be reproduced in any model, either in vitro or in silico. Animal studies offer an alternative but again ethical considerations and concerns are frequently raised regarding their relevance to human systems (Wickham et al., 2006).
2.4.2 In vitro models

*In vitro* models can be useful tools in developing structured foods that guarantee their safety in human subjects. In order to design foods with specific functions, *in vitro* gut models have been developed to study how food and drugs perform in the digestive tract. The importance of this is demonstrated by the wide variety of models that have been already developed. These models vary greatly in their complexity and type, as each has been developed for different applications.

GI tract *in vitro* models could be static or dynamic. Static or biochemical models such as membrane- and cell-based models are basically focused on studying the release, absorption and bioaccessibility of actives molecules. On the other hand, dynamic models simulate physical processing and physiological actions that occur *in vivo* such as shear rates, mixing or hydration, changing conditions over time (Kong and Singh, 2008a).

To study passive diffusion of actives through biological membranes and therefore predict the extent of absorption from the GI tract, membrane- and cellular- based models have been developed as explained below.

**2.4.2.1 Membrane-Based Models (PAMPA)**

This method provides an *in vitro* model for studying *passive diffusion*. As described in section 2.3.3, passive diffusion is an important transport mechanism in the GI tract to transport active molecules across cell membranes. The Parallel Artificial Membrane Permeation Assay (PAMPA) provides a straightforward approach to evaluate the permeability of various actives across artificial membranes. The use of artificial membranes to investigate permeation processes uses filter-supported lipid membrane systems to predict
the absorption profile of a drug, for example, where lipids composition can change accordingly. To make lipid membranes, phospholipids and other membrane constituents are added to the filter support as a solution in an organic solvent. As different tissues have different lipid composition; lipid composition is important and can be selected as required. Among the lipids available for designing these artificial membranes are: Phosphatidylcholine (PC), Phosphatidylethanolamine (PE), Phosphatidylinositol (PI) and cholesterol. Various pH profiles can be used to evaluate permeability (Deferme et al., 2008). They are widely used in pharma-related studies, providing information about the passive-transport permeability of a compound. The compounds can be categorised into low apparent permeability ($P_{app} < 10 \times 10^{-6}$ cm/s) and high ($P_{app} > 10 \times 10^{-6}$ cm/s) permeability (Deferme et al., 2008). The ability of this assay to evaluate permeability over a large pH range is valuable for an early understanding of how new oral compounds might be absorbed across the entire gastrointestinal tract.

2.4.2.2 Cell-based models

Cell based assays, such as Caco-2, assess permeability by transcellular passive diffusion as well as active and paracellular transport. Caco-2 cell systems are the most extensively characterized cell-based model because of their morphological and functional characteristics of the small intestinal absorptive cells, including the effective permeability (Deferme et al., 2008). The Caco-2 model is an *in vitro* static model consisting of one single cell type. One of the disadvantages of these models is that low molecular weight hydrophilic compounds showed poor permeability (Deferme et al., 2008). Caco-2 cells monolayers have been applied as an effective screening tool to address food factors associated with iron availability, deficiency and anemia, for example. The PAMPA and Caco-2 assays can be valuable when used in conjunction to identify the cause of poor absorption.
2.4.4.3 Mechanical models

Mechanical models have a wide selection features and different degrees of complexity depending on their intended final application. For instance, Arnold and Dubois (1983) simulated gastric mixing in a silicone rubber tube (i.d. 19 mm) placed in a peristaltic pump. The frequency of peristalsis was adjusted to 0, 1 and 3 waves per minute. The authors found that the diameter of solid food particles was reduced when the rate of contractions was increased. Food disintegration was also influenced by composition of fluid surrounding particles and reaction time.

De Boever et al. (2000) developed a model composed of a series of five double-jacketed vessels. This successive system was used as a Simulator of the Human Intestinal Microbial Ecosystem (SHIME) and was first developed by Molly et al. (1993). The original model consisted of five computer controlled reactors mimicking the conditions of duodenum/jejenum, ileum, caecum ascending, transverse and descending colon. De Boever et al. (2000) included a sixth reactor to mimic the chemical reactions occurring in the stomach. The SHIME is a validated model which provides more reproducible results for mechanistic studies. This model has been used to simulate the kinetics of the chyme passage through the GI tract; gastric enzymes, bile and small intestinal enzymes, pH, temperature, feed composition, transit time in the GI tract, anaerobic environment. However, absorption and feedback mechanism are not included.

Similar studies have been done by Probert and Gibson (2004), in which chemostats were used to study bacteria inside the intestine. The bioavailability of actives has been also tested in food and pharmaceutical areas using similar methods. For instance, the bioaccessibility of
ingested mycotoxins from food and soil contaminants have been studied by Versantvoort et al. (2005) and Oomen et al. (2003), respectively.

All mechanical models described above are mixed with an impeller, a magnetic stirrer, an orbital shaker or in the worst scenario are not mixed at all. However, the kneading type of mixing as occur in vivo cannot be simulated by any of these devices. As there is evidence that intestinal wall movements influence actives absorption, the next section is focus on discussing in vitro models that were implemented with mechanism of mixing with physiological relevant features, i.e. peristalsis and segmentation movements are mimicked by different mechanisms.

Macagno et al. (1982) developed a physical and mathematical model of the small intestine to study absorption by mimicking different types of contractions and therefore degrees of convective transport. The physical model used by these authors consisted of a dialysis tubing 1.6 cm in diameter connected to two reservoirs and immersed in a water bath at a constant temperature. Mixing contractions were produced by two rotating parallel cylinders mounted symmetrically. Two configurations were used to produce either a net flow in one direction or a back-and-forth flow when one end of the tube was closed. Oscillation flow was produced rather than through flow in order to simulate conditions more like those of the intestine. They measured the permeability of the tube wall when the tubing was filled with an aqueous solution of sodium chloride.

The mathematical model was developed from equations of conservation of volume, mass and momentum in which the wall movements, mass transfer and flow were taken into account. The effect of different type of contractions on diffusion was also evaluated. Results showed
that different types of contractions produced different degrees of convective transport. Absorption was increased by up to 30% when symmetric non-propagative contractions were induced, more than doubling (70%) due to asymmetric propagative contractions and increased almost 100% for symmetric propagative contractions, showing that wall movements enhanced mass transfer across the wall.

Edwards et al. (1988) demonstrated the importance of diffusion and convection of active molecules across the unstirred layer in viscous solutions by means of an in vitro model composed of dialysis tubing (12 x 1.5 cm). The tubing was tied at each end and anchored in a Perspex reservoir containing distilled water. Intestinal contractions were simulated using two pairs of paddles located at each end of the tubing. The volume of solution introduced into the tubing was less than its capacity but sufficient to be displaced backwards and forwards along the tubing by alternate contractions at either end of the dialysis tubing without causing distension of the tubing. The motor was set to generate either 36 or 72 contractions per minute. The effect of food formulation was also evaluated by adding 0.5% of guar gum to the solution of NaCl. They concluded that guar gum has no significant effect on diffusion of sodium. However, convection was greatly inhibited in the presence of guar gum. This biopolymer delays the mixing of solutions of differing conductivity. Reduction in absorption by guar gum is probably due to resistance of convection by viscous solutions of the convective effects of intestinal contractions.

Large and more sophisticated models include: (i) the TIM model from the TNO Nutrition and Food Research (Zeist, The Netherlands) and (ii) the Gut Model developed in the Institute of Food Research in Norwich, UK.
TNO in the Netherlands has developed two models: the Gastro Intestinal Tract Model (TIM-1) and the Large Intestinal Model (TIM-2). The TIM-1 is a dynamic in vitro model composed of four compartments to simulate the: stomach (A), duodenum (C), jejunum (E) and ileum (G), as illustrated in Figure 2.11. Each compartment is connected by two basic units and equipped with a flexible membrane inside, and outer glass jackets that allow the passage of water for both temperature and pressure control. The TIM mimics the interactions with gastrointestinal fluids and reactions while peristalsis is reproduced by altering external pressures; however, segmentation is not included in this model. Human physiological conditions in the stomach and the small intestine are mimicked, including simulation of pH changes, temperature, peristaltic movements, and secretion of digestion enzymes, bile, pancreatic juices, and absorption of active molecules (Kong and Singh, 2008a). The TIM has been used to validate the “biodrug” concept and to study the behaviour of orally administered drug dosage forms under physiological conditions (Blanquet et al., 2001). This model has been used in nutritional studies (Mateo Anson et al., 2010), to evaluate the bioavailability of active components (Duchateau and Klaffke, 2008, Minekus et al., 2005) and stability of antioxidants such as xanthophylls (Blanquet-Diot et al., 2009); and micronutrients such as iron, folic acid and ferulic acid (Mateo Anson et al., 2009). Studies indicate that the TIM shows to be a useful tool for various research purposes dealing with; availability for absorption of mutagenic; as well as antimutagenic components in food, probiotics and pharma studies to design and develop of new drug formulations and to predict the performance of several formulations in vivo.
In addition, the TIM-2 was designed under the same principles that the gastrointestinal model TIM-1, with flexible walls between which the intestinal content is circulated and a glass jacketed where water regulates the temperature and peristaltic movements of the chyme.

The Institute of Food Research (Norwich, UK) has recently developed a fully automated dynamic gastric model that simulates the human stomach digestion using a conical flexible walled vessel and a cylinder that processes the food at representative shear rates. This model simulates with accurate replication the physical mixing, transit and breakdown forces in the stomach, as well as the physiological conditions (pH gradient and enzymes) and emptying cycles measured within the main body of the human stomach (Marciani et al., 2001). The Model gut (Figure 2.12) can processes real food and drugs adjusting the residence time, acid...
and enzyme secretions according to the components of the food. It has been recently used to study the effect of allergens in foods (Chambers et al., 2004), release of nutrients or bioactive components (Vardakou et al., 2011), performance of functional foods and survival of probiotics (Lo Curto et al., 2011, Mandalari et al., 2010a).

Kong and Singh (2008b) also developed a model stomach system to investigate disintegration kinetics of solid foods during digestion. The model stomach consisted mainly of a turntable and a jacketed glass chamber containing simulated gastric juice in which plastic beads were added to simulate food particulates as well as to provide a suitable mechanical destructive force on food samples (Kong and Singh, 2008b). The disintegration of foods is simulated by applying representative shear stresses to break the food structure and evaluate the effect of microstructure on digestion. They reported that a good correlation was observed between the kinetics of food disintegration and in vivo stomach emptying.
The effect of wall motility particularly segmentation movements on absorption profiles was recently studied by a dynamic *in vitro* model of the small intestine at the University of Birmingham (UK). The model consists of a dialysis membrane and a mechanical system that simulates segmentation movements. The Small Intestine Model (SIM) allows the study of motility and chemical digestion including food digestion and actives absorption (Tharakan et al., 2010). The SIM was used for studying mass transfer, starch digestion and flow visualization of the lumen contents. While simulating the small intestinal flow profile, experiments have shown that the functional ingredient guar gum reduces the mass transfer coefficient by up to 30% of the model nutrient riboflavin. This together with computational modelling suggests an explanation for the observed functionality of guar gum to reduce the peak increase in blood glucose levels after ingestion of test meals. This is the starting point of this thesis.

### 2.4.3 *In silico* and theoretical models

Over the last decades, biological processes have been studied by developing mathematical models in order to obtain a fundamental understanding of the effect of a variety of physiological processes. *In silico* models have been used to evaluate the residence time distribution, release and inactivation of bioactives such as probiotics, drugs and foods in the GI tract mimicking the intestinal fluid flow (Stoll et al., 2000). These models have proved to be useful tools to predict the human fraction dose absorbed from orally administrated drugs.

Computational models have been created to study absorption processes in the GI tract. Although these models were fitted to the experimental data their use is limited due to the complexity of the problem. As a first approach, an equation based model without considering complex fluid dynamics must be developed. Computational Fluid Dynamics (CFD) has been
used to simulate segmentation and peristalsis in these *in vitro* and *in silico* models describing momentum transport during single phase flow which incorporates moving boundaries (Tharakan, 2008). Consideration was given to the design to ensure that the mechanical processing of the digesta in the human small intestine was being accurately represented.

Pal et al. (2004) for example, developed a two-dimensional computer model of the human stomach based on the “lattice-Boltzmann” method. The model demonstrated that antral contraction waves (ACW) are responsible for gastric mixing. Thus, the strongest fluid motions were found in the antrum with two basic flow patterns: (i) retropulsive flow velocities up to 7.5 mm/s and (ii) recirculating flow velocities of 2.0 mm/s. The authors concluded that both flow patterns enhance mixing process in complementary ways in the stomach. While particles are separated longitudinally by retropulsive flow, they are transported transversally to the gastric wall due to recirculation flow patterns.

Moreover, there are several commercially available software packages, e.g. GastroPlus™, iDEA™, Intellipharm PK and PK-Sim, all capable of reproducing and predict the human absorption processes with fairly high degree of accuracy. These based *in silico* models have been designed for the prediction of oral drug absorption (Johansson and Paterson, 2008).

On the other hand, theoretical and phenomenological models have been also designed to study, for instance, digestion and gustatory responsiveness for insect herbivores using chemical reactor models (Wolesensky et al., 2005).

Based on macrotransport analysis Stoll et al. (2000) developed a theoretical model for molecular absorption from the small intestine. The inputs used in the model were primarily
anatomical and physicochemical features from the literature. Complex convection, dispersion, degradation and absorption rates were included in this model. Furthermore, differential mass balance approaches have been developed to describe simultaneous fluid flow and drug absorption at a non-steady state of drug perfusion through the intestine as follows: dispersion model used to simulate oral drug absorption in which the change in concentration with distance and time is equal to the longitudinal spreading minus flow and absorption (Equation 2.1)

\[
\frac{dC(x,t)}{dt} = \alpha \left( \frac{d^2C(x,t)}{dx^2} \right) - \beta \left( \frac{dC(x,t)}{dx} \right) - k_a C(x,t)
\]

where \(C(x,t)\) is the drug concentration at any distance along the intestine at time \(t\) (mass/m\(^3\)), \(\alpha\) is the longitudinal spreading coefficient (m\(^2\)/s), \(\beta\) is the linear flow velocity (cm/s) and \(k_a\) is the apparent first-order absorption rate constant (per second).

As described above, a wide variety of in vitro and in silico models are available. Each of them developed with the same objective predicts the performance of an active ingredient in the GI tract.

2.5 The small intestine bioreactor: modeling digestion with chemical engineering

From an engineering perspective, the small intestine is a chemical bioreactor in which flow and mixing of chyme are crucial to enhance chemical digestive reactions and actives absorption. The simplest reactor configuration for an enzyme reaction is the batch reactor into which all reactants are initially loaded (Penry and Jumars, 1986, Levenspiel, 1999). An ideal batch reactor is assumed to be well mixed so the contents are uniform in composition at all times. On the other hand, in a plug-flow reactor or tubular flow reactor the substrate enters one end of a cylindrical tube. Since the long tube and lack of stirring device prevents complete mixing of fluid in the tube, the properties of the flowing stream will vary in both
longitudinal and radial directions (Levenspiel, 1999). Since the variation in the radial
direction is small compared to that in the longitudinal direction, it is called a plug-flow
reactor. Due to its geometry and flow characteristics, the small intestine resembles a plug
flow reactor in which lumen contents will also vary in both longitudinal and radial directions.
Fluid dynamic processes occurring in the human small intestine are a result of intestinal
motility, specifically segmentation movements (section 2.3.2), and are responsible for mixing
lumen contents. Therefore, intestinal motion primarily defines the fluid mixing profiles of the
food materials in the small intestine; mass transfer occurs through convective mixing and
diffusion processes taking place on a non-newtonian fluid (Tharakan, 2008). To understand
these processes is important to review some basic concepts that could be applied at digestion-
absorption phenomena in the small intestine.

2.5.1 Fluid dynamics

The study of fluid dynamics involves application of the fundamental principles of general
mechanics to liquids and gases. These principles involve conservation of matter, energy and
Newton’s laws of motion (Massey and Ward-Smith, 1998). Overall a fluid flow can be
divided into two types: laminar and turbulent flow. Laminar flow occurs when a fluid flows
in parallel layers without disruption between the layers. In turbulent flow, however, the fluid
undergoes irregular fluctuations and circulating vortices result in transference of fluid on a
larger scale, with cyclic fluctuations in the flow rate. Turbulent flow is a flow regime
characterized by property changes, low momentum diffusion and high momentum convection
and variation of pressure and velocity in space and time (Re > 4000).

Turbulent flow is often more desirable to many engineering process as it gives greater mass
transfer (Cussler, 2009). Particles of fluid which are initially separated by a long distance can
be closed together by eddy motions in turbulent flows. As a consequence, heat, mass and momentum are very effectively exchanged. Such effective mixing gives a rise of diffusion coefficients for mass, momentum and heat transfer.

The type of flow is dependent on the velocity, physical properties of the fluid and the geometry of the surface (Coulson and Richardson, 1998). To describe these properties a suitable correlation widely used by engineers is the Reynolds number (Re) which gives the relation between inertia forces and viscosity forces; for high Re numbers inertia is greater than viscous forces and the flow is turbulent, when the Reynolds number is low, viscous forces are greater and the flow is laminar. Re can be calculated using Equation 2.2.

\[
Re = \frac{\rho v D}{\mu}
\]

(2.2)

where \(\rho\) is the density of the fluid, \(v\) the mean velocity, \(D\) is the diameter of the pipe and \(\mu\) the viscosity of the fluid. For pipe flow, laminar flow occurs when \(Re < 2100\), while turbulent flow occurs when \(Re > 10000\).

In digesta, solid particles can also modify the flow properties, but by comparison with soluble polysaccharides, their intrinsic viscosity is lower. The overall viscosity of fluid digesta (stomach and small intestine) can be increased by significant amounts of soluble polysaccharides. Evaluating the viscosity generated by soluble dietary fibres in dilute digesta is not an easy task, also because their concentration in the lumen is different from those ingested, as shown by Marciani et al. (2000). In particular, the total volume of digesta may adapt in response of ingestion of viscous solution, partly offsetting the difference in initial viscosity (Guillon and Champ, 2000).
Fluid flow in the small intestine is provided by the peristaltic movements that bring chyme from stomach to the small intestine. The velocity of peristaltic flow as well as the physical properties of the chyme will determine the flow type either laminar or turbulent.

As described in section 2.3, the small intestine model can be considered as a baffled reactor due to the intestinal wall structure in which oscillatory Reynolds numbers describe the type of flow occurring in the small intestine, enhancing heat and mass transfer processes. Oscillatory Reynolds numbers can be calculated by Equation 2.3.

\[
\text{Re}_o = \frac{\rho 2\pi f x_o D}{\mu}
\]  

(2.3)

where \( \rho \) is the fluid density, \( f \) the frequency of oscillation, \( x_o \) is the centre-to-peak amplitude, \( D \) is the tube diameter, and \( \mu \) the viscosity of the fluid.

While there is some information from animal models, shear rates and flow profiles have not been well established for the human small intestine due to its complexity and location (Tharakan, 2008). For instance, Jeffrey et al. (2003) modelled flow profiles numerically, using inputs from luminal pressures, outflow and wall movements of isolated segments of guinea pig ileum.

### 2.5.2 Mixing fundamentals

Mixing is one of the most common operations in chemical and biochemical processing. It is used to enhance mass transfer processes, dissolving the solute homogenously in the medium and reducing the gradient of concentration. Mixing is provided by the segmentation movements of the intestinal wall while motility in the gastric antrum plays an important role in mixing and food transport. Mechanical agitation in the gut promotes not only chemical
reactions but also mass transfer processes. Mixing by eddy diffusion is much faster than mixing by molecular diffusion and, consequently, turbulent mixing occurs much more rapidly than laminar mixing.

Mixing mechanisms (Paul et al., 2004):

- **Molecular diffusion** is diffusion caused by relative molecular motion and is characterized by the molecular diffusivity $D_{AB}$. Diffusion is responsible for the fluid micromixing at small scales.

- **Eddy diffusion** or **turbulent diffusion** is dispersion in turbulent flows caused by the motion of large groups of molecules known as eddies. This motion is measured as the turbulent velocity fluctuations. The turbulent diffusivity, $D_t$ is a conceptual analogy to $D_{AB}$ but is a property of the local flow rather than the fluid.

- **Convection** is dispersion caused by bulk motion at large length scales, driven by gross flow or macromixing.

- **Taylor dispersion** is a special case of convection, where the dispersion is produced by velocity gradients across the flow. It is most often referred to in the case of laminar pipe flow, where axial dispersion arises due to the parabolic velocity gradient in the pipe, i.e. the centre of the flow is moving faster than that at the walls.

In a flowing fluid, the rates of transfer processes and reactions are affected by the flow. Convective mixing of the chyme has been recognized as one of the functions accomplished by intestinal motility, but besides general statements to this effect, little is known about the quantitative nature of such mixing. A first step into gaining some insight into the transport of mixing of the chyme is achieved if a proper visualization of the flow induced by the wall movements is attained (Macagno and Christensen, 1980).
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2.5.3 Molecular delivery

The delivery of active molecules to the intestinal wall is a result of both diffusion and convective mixing of the intestinal contents (digesta). Molecular delivery occurs by convection and diffusion across the intestinal wall or membrane. Mass transfer could be described by two models, represented by using either diffusion or mass transfer coefficient.

The concentration gradient between the fluid inside the small intestine and blood outside drives the diffusion phenomenon (Tharakan, 2008). Diffusion and convection are briefly discussed in the next section.

2.5.3.1 Diffusion and convection

Diffusion is a process by which molecules, ions, or other small particles spontaneously mix, moving from regions of relatively high concentration into regions of lower concentration. The rate of movement of material by diffusion is described mathematically by Fick’s laws. Fick’s first law is described by Equation 2.4, which relates the diffusion flux \( J \) to the gradient of the change in concentration \( (dc/dx) \).

\[
J = -D \frac{dc}{dx} \tag{2.4}
\]

where \( J \) is the diffusion flux of material per unit area (mol/m\(^2\)/s); measures the amount of substance that will flow through a small area during a small time interval, \( D \) is the diffusion coefficient or diffusivity in (m\(^2\)/s), \( c \) (for ideal mixtures) is the concentration (mol/m\(^3\)) and \( x \) is the position (m). The negative sign indicates that flow is down the concentration gradient (from high to low concentration).

\( D \) is proportional to the squared velocity of the diffusing particles, which depends on the temperature, viscosity of the fluid and the size of the particles according to the Stokes-
Einstein relation. In dilute aqueous solutions the diffusion coefficients of most ions are similar and have values that at room temperature are in the range of $0.6 \times 10^{-9}$ to $2 \times 10^{-9}$ m$^2$/s. For biological molecules the diffusion coefficients normally range from $10^{-11}$ to $10^{-10}$ m$^2$/s.

Fick's second law for unidimensional flow (Equation 2.5) describes how concentration changes as a function of time. The rate of change of the concentration as a function of time is a function of the change in concentration gradient (Cussler, 1997).

\[ \frac{\partial c}{\partial t} = D \frac{\partial^2 c}{\partial x^2} \quad (2.5) \]

where $c$ is the concentration (mol/m$^3$), $t$ is time (s), $D$ is the diffusion coefficient (m$^2$/s) and $x$ is the position (m).

On the other hand, convective transport occurs when a constituent of the fluid (mass, energy, a component in a mixture) is carried along with the fluid. Convective heat and mass transfer take place through both diffusion random Brownian motion of individual particles in the fluid, and by advection, in which matter is transported by the larger-scale motion of currents in the fluid. By the nature of the small intestine, convention is a result of induced flow by the intestinal movements that transport and mix chyme along the intestine ‘force convection’.

2.5.3.2 Mass transfer coefficients

Assuming that the lumen content is a well-mixed solution, the mass transferred from some interface is proportional to the concentration difference and the interfacial area as stated in Equation 2.6 (Cussler, 2009).
(amount of mass transferred) = K(interfacial area)(concentration difference) \hspace{1cm} (2.6)

where $K$ is the mass transfer coefficient (m/s).

Dividing both sides of the equation by the area, equation (2.6) could be arranged as:

$N_i = k(c_{li} - c)$ \hspace{1cm} (2.7)

where $N_i$ is the flux at the interface and $c_{li}$ and $c$ are the concentrations at the interface and in the bulk solution, respectively.

The simplest theory for interfacial mass transfer, assumes that a stagnant film exists near every interface. This film, also called an unstirred layer, is almost always hypothetical, for fluid motions commonly occur right up to even a solid interface. The steady-state flux across this thin film can be written in terms of the mass transfer coefficient.

A common dimensionless number used in the analysis of mass transfer phenomena is the Sherwood number ($Sh$) or mass transfer Nusselt number ($Nu$) that represents the ratio of convective to diffusive mass transport (Equation 2.8).

$$Sh = \frac{k_b p L}{D_{AB}} = \frac{\text{Convective mass transfer coefficient}}{\text{Diffusive mass transfer coefficient}} \hspace{1cm} (2.8)$$

where $L$ is a characteristic length (m), $D_{AB}$ is mass diffusivity (m$^2$/s) and $k_b p$ is the local mass transfer coefficient (m/s).

The flow within a given process will be highly complex and difficult to evaluate fully and even if a full CFD model were available, it is likely that it would be too complex to use in practice. Mass transfer coefficients are defined in terms of measurable quantities, such as the bulk concentration, the concentration in the receiving medium, and the flux of mass between
the two. The active concentration is assumed to have an average value, and the flux between the two regions is related to the difference between these average concentrations. Thus, mass transfer coefficients are used to describe how the concentration changes with position or how concentration changes with time (Cussler, 2009).

The convective mixing of intestinal contents is one of the functions of intestinal motility. In practice little is known about the quantitative nature of this mixing. Mass transfer experiments, visualisation and modelling of the flow, induced mixing by the wall movements, will achieve an insight into the transport and mixing of digesta and how food formulations can affect this (Tharakan, 2008).

In order to study flow as a function of the intestinal motility, techniques such as Planar Laser-Induced Fluorescence (PLIF) and Particle Image Velocimetry (PIV) have been successfully used in problems of fluid flow visualization in which transparent solutions are used (Hu et al., 2012, Li et al., 2011). Both are powerful flow diagnostic techniques based on image analysis. Whilst PLIF is widely used to monitor macromixing time, PIV is commonly employed to study velocity distributions. PLIF is a non-invasive technique in which a tracer e.g. rhodamine, is injected into flow which fluoresces upon excitation by laser light. PLIF supplies conventional two-dimensional images which are digitally processed. The greyscale intensity of the images can be related to local tracer concentration. Spatial resolution however is limited by the camera magnification; image can be processed pixel by pixel is required. In PIV paired laser pulses illuminate neutrally buoyant seeding particles suspended in flow. Here, statistical analysis of particle images yields information regarding local fluid velocity.
However, one of the problems is that the above techniques need transparent fluids, i.e. their use is limited. A very powerful alternative technique is the Positron Emission Particle Tracking (PEPT). PEPT is a variant of the Positron Emission Tomography (PET), which is well established in medicine for imaging the human body. In terms of engineering applications, the PEPT technique has been utilised to study solids motion within a variety of equipment including spouted beds, rotating cans fluidised beds and mixers (Cox et al., 2003, Kuo et al., 2002, Seiler, 2001).

### 2.6 Food formulation (structured foods)

Intestinal absorption is not only related to the flow and mixing of fluids in the gut but also to the structure of the food within the intestine. It has been well established that food structure has a significant impact on the food digestive behavior (Lundin et al., 2008). For instance, it has been demonstrated that by adding dietary fibre such as guar gum, pectins, β-glucans, cellulose and fibres into foods can notably delay absorption of glucose into the bloodstream (Blackburn et al., 1984, Jenkins et al., 1978, Salmerón et al., 1997).

Dietary fibre is defined as a heterogeneous mixture of substances predominantly found in plant cell wall that are not hydrolised by the digestive enzymes in man. Indigestibility implies that dietary fibre is not broken down by enzymes in the upper part of the gastrointestinal tract, because the human small intestine does not contain enzymes capable of cleaving β-1-4 bonds present in dietary fibres. Dietary fibre or Non Starch Polysaccharides (NSP) are used as a ‘fuel’ for the gut microbiota which promotes the production of short fatty acids in the large bowel that are easily absorbed and contribute to energy metabolism.
Dietary fibre is classified into two groups, water-soluble and water-insoluble fibres (Anttila et al., 2004). Soluble viscous fibres contribute to formations of the unstirred layer adjacent to the mucosa, which serves as a physical barrier to nutrient absorption and bile acid reabsorption. Water insoluble fractions; such as cellulose and lignin and water-soluble components, encompassing the major part of hemi-cellulose compounds, gums, β-glucans, mucilages and pectins (Ryttig, 2005).

The two major benefits associated with soluble and viscous dietary fibres are attenuation of glycemic response and plasma cholesterol lowering (Anttila et al., 2004). It is generally recognized today that the physiological value of soluble dietary fibre is based on its ability to increase the viscosity of food digesta in the intestine (Anttila et al., 2004). Studies have shown that addition of soluble fibres such as pectin, guar gum and locust bean gum in food reduces gastric emptying, delays absorption of actives, reduces the plasma glucose response and slows down the return of hunger. Soluble fibres have been combined in diet for treating pathological conditions such as obesity, hypercholesterolemia and diabetes (Kong and Singh, 2008a).

The rate of starch hydrolysis and glucose absorption could be reduced when foods are supplemented by gelling fibres such as guar gum, pectin, tragacanth or cellulose. It has been stated that postprandial glucose levels have been reduced when foods are supplemented with fibres. Gastric emptying and intestinal motility are delayed or modified when viscous biopolymers are added to foods, slowing glucose diffusion through the blood stream because of increasing of the unstirred layer from the chyme to the epithelium of the small intestine. Furthermore, amylase accessibility is reduced as a result of increased viscosity of lumen.
contents (Leclere et al., 1994, Snow and O'Dea, 1981). Fibre could “hide” or isolate starch from amylases avoiding formation of the complex starch-enzyme delaying starch hydrolysis.

Addition of guar to nutrient drinks and solid meals markedly prolongs the duration of a postprandial motility pattern (Schönfeld et al., 1997). Meyer et al (1988), suggested that guar slows emptying of the liquid part of the meal, but speeds emptying of particulate matter. Gastric sieving may be impaired by guar (Meyer et al., 1986 and 1988).

Blackburn et al. (1984) found that by adding 9 g of guar to a liquid model food containing glucose significantly improve glucose tolerance; as one can see in Figure 2.13, compared with the control, blood glucose levels were lower at 30 and 40 minutes. These authors also observed a delay in gastric emptying of the glucose drink when guar was added to the solution.

![Figure 2.13](image-url)

**Figure 2.13** Blood glucose levels of human subjects following ingestion of 50g glucose in 250ml orange drink, with or without 9g guar gum. Results are expressed as a mean with the standard errors of six observations (taken from Blackburn et al. 1984).
The rate of intestinal digestion, absorption and transit of the food could be also influenced by the physical form of food ingested and the physical properties of the intestinal contents produced (Jenkins et al., 1982).

Overall, the food industry needs to develop foods that deliver active ingredients with specific functions over an extended period of time. Potentially, this could be accomplished by changing food formulation and developing encapsulated structures which can be released at the right place and time. Two approaches are attempted in this thesis:

(i) The change in bulk viscosity of fluid in a model gut and
(ii) A model encapsulation system

By changing the bulk viscosity of the intestinal fluid, the biopolymers will interact with each other because of entanglements i.e. no permanent bonds between individual polymers are formed. On the contrary, in the case of polysaccharide gels (networks) or microstructures formation, the polymers are cross-linked into a tridimensional structure and the addition of solvent, in most cases, not change the integrity of the network. In both cases, a physical barrier is developed to delay the release, digestion and further absorption of key molecules to design functional and structured foods with health benefits to the consumer.

Therefore, a briefly review about encapsulation, wall materials and techniques such as ionotropic gelation and liposomes is presented below.

### 2.7 Encapsulation

Encapsulation is a physicochemical process widely used in different areas such as pharmaceutical, textile, agrochemical and foods, aiming to protect a bioactive ingredient against a variety of environmental conditions (e.g. pH, temperature, oxygen).
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Microencapsulation has been used to pack solids, liquids or gaseous materials into micro- or nano-size sealed capsules for release at controlled rates using desired released triggers (Klaypradit and Huang, 2008, Pothakamury and Barbosa-Cánovas, 1995).

The compounds that have been more commonly encapsulated in the food industry include: proteins e.g. insulin; probiotics (Annan et al., 2008), (Soottitantawat et al., 2005), antioxidants and nutraceuticals (Chiou and Langrish, 2007), vitamins (Chen and Wagner, 2004), minerals (Gouin, 2004), essential oils (Ahn et al., 2008) and sugars like sucrose encapsulated in oil cotton (Jozwiakowski et al., 1990). The simplest microcapsulate consists of a core or active ingredient surrounded by a wall, shell or barrier. Nevertheless, whether the active ingredient is embedded inside of a matrix the structure is known as a microsphere. The core is the component that required protection and it may be composed of one or more ingredients (Klaypradit and Huang, 2008). The wall could be single or multilayer (Pothakamury and Barbosa-Cánovas, 1995). Encapsulation techniques used in this thesis include

2.7.1 Ionotropic gelation

Ionic gelation is one of the methods that much research attention has attracted, to the formation of microspheres using macromolecules that can bond divalent ions such as Ca $^{++}$. It has been used for insulin-chitosan particles nanoencapsulation, adding a tripolyphosphate (TPP) solution (Agnihotri et al., 2004). Moreover, different wall materials have been used such as pectin and alginate with calcium chloride as a cross-linking agent, obtaining good results in the entrapment of drugs and foods (Goh et al., 2012, Annan et al., 2008, de Vos et al., 2006, Draget et al., 1994).
2.7. 2. Liposome entrapment

It is a process commonly used by the pharmaceutical industry for the protection of vaccines, enzymes or drugs (Feng et al., 2004a). The capsules produced by this technique are formed by a phospholipid, such as phosphatidylcholine, which may include in its structure hydrophilic or hydrophobic material providing greater versatility and endurance (Lee et al., 2008, Colletier et al., 2002, Kokkona et al., 2000, Kim and Baianu, 1991).

The most common shell materials used in the food industry are listed in Table 2.1.

Table 2.1 Common materials used in food microencapsulation

<table>
<thead>
<tr>
<th>Component</th>
<th>Examples</th>
<th>References</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Carbohydrates</strong></td>
<td><strong>Starch derivates:</strong> Modified resistant starches, Cyclodextrins, Maltodextrins (DE &lt; 20), Corn syrup solids (DE &gt; 20)</td>
<td>Dimantov et al. (2004)</td>
</tr>
<tr>
<td></td>
<td><strong>Cellulose derivates:</strong> Carboxymethyl-cellulose (CMC), Hydroxypropylmethyl-cellulose (HPMC)</td>
<td>Shahidi and Han (1993)</td>
</tr>
<tr>
<td></td>
<td><strong>Hydrocolloids:</strong> Arabic gum, mesquite, Carrageenan, Pectin, agar, Chitosan, Sodium alginate</td>
<td>Su et al. (2008), Krishnan et al. (2005), Shahidi and Han (1993), Drush, (2007), Agnihotri et al. (2004), Gouin (2004)</td>
</tr>
<tr>
<td><strong>Proteins</strong></td>
<td>Gelatin, Milk proteins: casein, albumins, Wheat, corn and soy proteins</td>
<td>Shadini and Han (1993), Je Lee and Rosenberg (2000), Yoshii et al. (2001)</td>
</tr>
<tr>
<td><strong>Lipids</strong></td>
<td>Wax, paraffins, Mono-, di- glicerides, Fosfoglicerides, Oils, fats</td>
<td>Shadini and Han (1993), Feng et al. (2004b)</td>
</tr>
</tbody>
</table>

*DE = Dextrose Equivalent*
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The wall materials used in the encapsulation of an active substance must have emulsification and film forming properties, give low viscosities at high solids concentration (300 cps > 35% solids), have low hygroscopicity, bland taste and low-cost, release the active ingredient when it is reconstituted, provide good protection against environmental conditions (Shahidi and Han, 1993). Some of the materials above could be employed in combination with others to improve their film forming properties, for instance, it have been used successfully starch - gum arabic or chitosan – alginate (Agnihotri et al., 2004) for encapsulating agents flavoring mixtures. In addition, the pharmaceutical industry uses enteric materials such as “Eudragit®”, to encapsulate different types of drugs. These enteric coatings have different dissolution profiles according to the pH and are designed to release the active ingredient at a specific pH. Most of these materials are methacrylate polymers able to protect the active material against the acidic pH of the stomach.

2.8 Conclusion

Despite of its importance on the delivery of active molecules relatively little work has been done to understand and study mass transfer phenomena inside the small intestine. In summary the study of physiological processes involved in food digestion and active absorption is quite challenging, offering an infinite number of possibilities and opportunities in the design and develop of structured foods.

The first section of this Chapter covered the basic aspects of the GI tract including: anatomical features, physiology, mechanical and chemical digestion, together with absorption and some relevant models developed to study these phenomena in vitro. According to this information there are opportunities to design and develop structured foods with health benefits for the consumers.
CHAPTER 3 - MATERIALS AND METHODS

3.1. Introduction

As discussed in the previous chapters, the design and development of novel functional foods require a better understanding of the impact that food formulation and structure have on digestion and absorption processes in the gastrointestinal tract. To predict the rate at which molecules are digested and absorbed through the epithelium, interactions between macronutrients after food processing must be also understood. The study of these phenomena ‘in vivo’ is very complex, expensive and time consuming. As such, realistic in vitro models must be used to give representative data of what happens inside the body during digestion. In this context, a dynamic ‘in vitro’ small intestine model (SIM) was used here to study the effect of mixing and formulation on mass transfer phenomena during nutrient digestion and actives absorption. The strategies proposed in this research were based on key features of the human small intestine taking into account its physiology and the physicochemical properties of the digesta.

This Chapter describes the materials and experimental procedures that were used to study mass transfer in the small intestine during digestion. As mentioned in the following sections, the effect of mixing and food formulation on glucose absorption was evaluated in the SIM under physiological conditions. Food formulation was varied by adding different concentrations of viscous fibres such as guar gum, pectin and cellulose. The second section of this chapter is focused on the hydrolysis of starch, since it is the main carbohydrate in human nutrition and its digestion in vivo is an indicator of the resulting postprandial glucose levels (Lehmann and Robin, 2007). Visualization techniques such as Planar Laser Induced Fluorescence (PLIF) and Positron Emission Particle Tracking (PEPT) were employed to
study velocity profiles and mixing performance in the SIM as a function of food formulation - viscosity. In the last section, a description of the encapsulation techniques used to produce structures for delaying the release of actives was included along with studies of entrapment efficiency, morphological characterization and in vitro release kinetics.

3.2 Materials

This section gives a brief description of all chemicals and enzymes used through this work, including: **monosaccharides and polysaccharides:** glucose and starch, respectively; **biopolymers:** guar gum, pectin, alginate, carboxymethyl cellulose and **enzymes:** α-amylase and amyloglucosidase for starch and food digestion. All reagents and digestive enzymes were ACS reagents purchased from Sigma-Aldrich Company (Poole, UK), unless otherwise specified.

3.2.1 Glucose (C$_6$H$_{12}$O$_6$)

Due to its importance as a biological fuel for most of the living cells, D-(+)-glucose (G5767 from Sigma-Aldrich, UK), was selected to model absorption in the small intestine. As an energy source, glucose plays an important role in the glycolysis metabolic pathway and it is critical in the synthesis of proteins, lipids and nucleotides. This monosaccharide is the basic unit of complex molecules such as starch, cellulose and glycogen. Glucose is the major sugar in blood and is constantly regulated by insulin. Insulin is a hormone, produced by the pancreas, which is essential in carbohydrate and fat metabolism regulation in the body. Insulin is responsible for cells in the liver, muscle, and fat tissue to incorporate glucose from the blood, storing it as glycogen inside these tissues (Guyton and Hall 2006). Normal blood glucose levels are between 4.4 mM and 6.7 mM in an adult. To low or high glucose levels in
blood can result in a health detriment. **Physical properties**: molecular weight 180.16 g/mol and melting point 150-152 °C.

### 3.2.2 Starch (C\(_6\)H\(_{10}\)O\(_5\))\(_n\)

Starch is a highly organized biopolymer composed of two polysaccharides: amylose (25%) and amylopectin (70-75%) arranged in starch granules. **Amylose** is a lower-molecular-weight (MW > 10⁵) crystalline linear biopolymer compressed by glucose residues linked by \(\alpha(1\rightarrow4)\) bonds (Figure 3.1a). On the other hand, the semicrystalline branched biopolymer **amylopectin** is a high-molecular-weight (MW > 10⁶), with a linear chain of \(\alpha(1\rightarrow4)\) glucose residues and branched point linked by \(\alpha(1\rightarrow6)\) bonds occurring every 25 units of glucose (Figure 3.1b). As starchy foods are very important part of our diet, starch from corn (S4126, Sigma-Aldrich, UK) was used to study digestion.

**Figure 3.1** Starch structure (a) amylose and (b) amylopectin (Chaplin, 2012)

### 3.2.3 Biopolymers and soluble fibres

Since soluble viscous polysaccharides such as guar gum, pectin and CMC have a significant effect on glucose and insulin tolerance when added to food (Ellis et al., 1995; Blackburn, Holgate and Read, 1984; Jenkins et al., 1978), they were chosen to study the effect of food
Chapter 3 - Materials and Methods

formulation on absorption by increasing the viscosity of the sample. As intestinal enzymes cannot hydrolyze these biopolymers, soluble fibres remain undigested in the small intestine.

### 3.2.3.1 Guar Gum

Guar gum is a galactomannan extracted from the endosperm of the seed of *Cyamopsis tetragonolobus*. The principal backbone is a chain of α-(1,4)-D-mannopyranosyl units with single α-(1-6)-D-galactopyranosyl units linked to the main chain. The mannose:galactose ratio is 1.2 to 1.8, however, this can change according to the temperature of the solution (Casas et al., 2000). Guar gum (105008, ICN Biomedicals, USA) has been employed in this research to modify the formulation of the food by improving viscosity, as it has shown to be one of the most effective soluble fibres to reduce the glucose blood levels (Blackburn et al., 1984).

### 3.2.3.2 Carboxymethyl cellulose (CMC)

Carboxymethyl cellulose sodium salt high viscosity (C5013, supplied by Sigma-Aldrich, UK) was used in glucose absorption studies to improve the viscosity of the sample. CMC is a slightly yellowish powder derived from cellulose with a linear chain of β-1,4 linked glucose units and carboxymethyl groups. Due to its properties CMC is used as a thickener, stabilizer and suspending agent. Like guar, by adding CMC the viscosity of the digesta can be modified.

### 3.2.3.3 Citrus pectin

Pectin is a poly-D-galacturonic acid methyl ester commonly extracted from the apple pomace and citrus peel. Due to its ability to form hydrogels upon addition of divalent cations such as Ca^{++}, pectin has been widely used in the food and pharma industries to produce gel
Chapter 3 - Materials and Methods

microstructures. Pectin could also be used as a functional ingredient in food or drink preparations to reduce postprandial glucose levels in both normal and diabetic patients. On this context, citrus pectin high methoxyl (7680, degree of esterification \textasciitilde 60\% from Fluka, UK) was used to increase the viscosity of glucose solutions and to estimate the effect that mixing frequency has on absorption. Compare to low methoxyl-pectin, high-methoxyl pectin was selected in this research due to its greater inhibitive effect on intestinal absorption of glucose (Kim, 2005).

3.2.3.4 Sodium Alginate

Extracted from the native brown seaweed \textit{(Phaeophyceae)}, alginate is an unbranched block copolymer composed of 1-4-linked residues of \(\beta\)-D-mannuronic acid and \(\alpha\)-L-guluronic acid. Since alginate produces hydrogel membranes for encapsulation, whose characteristics (gel thickness, volume, mechanical resistance and porosity) can be easily modified, encapsulation of actives was done by using sodium alginate as encapsulating material. Gelling microspheres were produced with sodium alginate supplied by Sigma-Aldrich, UK (section 7.2) whereas liposomes were made with sodium alginate isolated from laminaria \textit{Hyperborea}, high in guluronate residues, from FMC Biopolymer (Norway), section 7.3.

3.2.4 Enzymes

For starch digestion studies, two enzymes were used: \(\alpha\)-amylase isolated from porcine pancreas (EC 3.2.1.1. A3176 Type VI-B, \(\geq 30 \) units/mg solid, Sigma, UK) and amylglucosidase isolated from \textit{Aspergillus niger} (EC 3.2.1.3. A7095, \(\geq 300 \) U/mL, aqueous solution, Sigma, UK). Both enzymes were used together at ‘physiological concentrations’ to mimic the starch-containing food digestion in the intestinal lumen. Although data in the literature are sparse and hence difficult to define concentrations of pancreatic enzymes in the
Chapter 3 - Materials and Methods

duodenum, α-amylase secretions in adults has been reported between 5-15 nM (Slaughter et al., 2001). α-Amylase is a glycoprotein with a single polypeptide chain of ~475 residues containing two SH groups and four disulfide bridges. For activity and stability this enzyme requires both calcium and chlorine ions. As specified by the supplier the pH range for activity is from 5.5 to 8.0, with the pH optimum at 7.

Amyloglucosidase (AMG) from A. niger was used to convert maltose and maltodextrins to glucose in the lumen side of the SIM, mimicking hydrolysis occurring by the microflora present in the intestinal wall. AMG from Aspergillus or Rhizopus have been extensively used in mechanism of digestion in vitro to complete hydrolysis of carbohydrates to glucose (Englysh et al. (1992). Commercial pancreatin was not used in this work as the amylase activity in unknown and only starch was used as food model.

3.3. Small Intestine Model description

The small intestine model (SIM) used in this work, was designed and implemented by Tharakan (2008) in the School of Chemical Engineering at the University of Birmingham, UK. The SIM functions as a concentrically mass transfer exchanger composed of an inner semi-permeable RC membrane (Spectra/Por 7®, MWCO 8000 Daltons, 32 mm of diameter, Medicell International Ltd London, UK) and an outer non-active and impermeable silicone transparent tubing (Flexible Hose Supplies, UK, diameter of 50 mm and wall thickness 3 mm). This dynamic in vitro model has a unique pneumatic mechanism to reproduce the characteristic segmentation movements responsible for the flow and mixing in the human small intestine.
3.3.1. Membrane description/characteristics

A semi-permeable membrane of regenerated cellulose RC (Spectra/Por® 7 Dialysis Membrane Medicell International Ltd., London, UK) was used to represent the lumen side of the small intestine model (SIM). This is a flexible, transparent membrane with an approximate Molecular Weight Cut Off (MWCO) of 8000 Daltons, a nominal thickness of 0.05 mm and flat width of 50 mm. RC membranes are stable in a pH range of 5 - 9, compatible with most organics, with no fixed charges and no absorption of most of the solutes.

Although semi-permeable membranes have been widely used to mimic absorption of actives (e.g. TIM models), they have limitations on reproducing absorption processes. Within most of commercial membranes, only mechanisms of diffusion can be modelled. Their anatomical and physiological features restrict their used when active transport process of substances such as glucose, needs to be mimicked. A permeable membrane of cellulose has neither the characteristic anatomical structure of the intestinal wall (villus and microvilli) nor the plasma membrane of the epithelium cells as shown in Figures 2.9 and 2.10. Therefore, the area available for absorption in the SIM is reduced to 0.05 m² and re-circulation of intestinal content must be done. By using the SIM model only part of the absorption process can be predicted. On the other hand, the concentration of glucose in the recipient side of the SIM can be compared with blood glucose levels in adults (4.4 mM and 6.7 mM), after clearance of absorbed molecules from the circulation by the liver and the other organs and tissues.

3.3.2 The core section of the SIM: lumen and recipient sides

The SIM comprises by two independent and flexible compartments: the food or lumen side where digestion reactions take place and the recipient side where the products of absorption
can be collected. The inner side or lumen of the SIM is represented by the semi-permeable dialysis membrane described in section 3.3.1, with a characteristic length of 50 cm and 3.0 cm of diameter, whereas the outer or recipient side is represented by an outer transparent tube (50 cm and a 5 cm of diameter). Each side of the SIM, including the membrane and the transparent tube were holding on independent stainless steel tube claps with fluid inlets/outlets for the inner and outer side as shown in Figure 3.2a. The membrane and the tubing were held in place using on each end two rubber gaskets and two aluminum hose clamps, respectively, to form water tight seals and to avoid leaks during experimentation. The design used a length of 50 cm of tubing available for mass transfer. The flexible tubing allowed being deformed in a physiologically representative manner (Tharakan, 2008).

As shown in Figure 3.2 and 3.3, each compartment (lumen and recipient side) includes a peristaltic pump (Watson-Marlow, 323S/D) to feed and propels the solution into the system along with a surge tank or reservoir for fluids storage. The following lists the main components of the small intestine model (SIM) including the segmentation control section that will be described in detail in section 3.3.3.

### 3.3.3 The mixing or segmentation mechanism

Segmentation contractions in the SIM were produced by inflation and deflation of two inflatable rubber cuffs (from an Omron Blood Pressure monitor) around the wall of the intestinal rig (Figure 3.2a). When the cuff was fully inflated the external wall of the SIM was compressed and occluded, generating mixing and propulsion of the lumen content. On the contrary, when the cuff was deflated relaxation of the membrane was produced. In the SIM, inflation was induced by supplying pressurized air into the cuff whereas deflation was attained by applying vacuum. The air flow and vacuum were controlled by solenoid valves.
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(Burkert W2YLE, Germany) fixed on a control panel (Figure 3.2b). The opening or closing of the solenoid valves was automatically controlled by a programmable logic controller (PLC, National Instruments NI USB-9162) connected to a computer by the National Instruments NI-DAWMAX software. The PLC was powered by a 24V power supply (Power-one, 24VDC at 2.4A). A pressure indicator (Sunx: DP2-42E) was used to monitor the rate at which the cuff was inflated by the compressed air. Additionally, pressure relief valves (Norgen: 61B2) were implemented to the compressed air lines for safety reasons.

![Figure 3.2 Small Intestine Model (SIM), (a) membrane and inflatable cuffs and (b) solenoid valves panel to control the segmentation pneumatic mechanisms.](image-url)
Chapter 3 - Materials and Methods

I. Intestinal section

1. Peristaltic pump for inner (food) side (P1)
2. Peristaltic pump for outer (recipient) side (P2)
3. Feed and surge tank for the food side (T1)
4. Small intestine section outlet and feed tank inlet (food side)
5. Feed and surge tank for the recipient side (T2)
6. Recipient side section outlet and feed tank inlet
7. Inflatable cuffs (Omron) to replicate the segmentation action (C1, C2)

II. Segmentation control

1. Inlet pressure sensor connected to compressed air inlet line (Pi)
2. Pressure relief valves (PRV1, PRV2)
3. Solenoid valves (S1, S2, S3, S4)
4. Connections for solenoid valves to programmable logic controller (PLC) for open and closed settings. PLC controlled by settable software on PC
5. Connections to gas lines (compressed/vacuum)

![Figure 3.3 A schematic representation of the main components of the SIM.](image-url)
Chapter 3 - Materials and Methods

The SIM was designed to give flexibility on the segmentation sequences by controlling the inflation, deflation and delay time intervals (1-6 s) through a LabVIEW programme designed by Derek Green from the School of Biosciences in the University of Birmingham, UK.

3.4 Methods

This section is divided into four parts that explain in detail the methodology used to gain a better understanding of the impact that mixing and food formulation have on digestion and absorption occurring in the upper small intestine. The methods described here are organized according to the next chapters as follows:

(i) Absorption of actives (Chapter 4)
(ii) Starch digestion and glucose absorption (Chapter 5)
(iii) Flow visualization techniques (Chapter 6) and
(iv) Encapsulation of actives (Chapter 7)

Experiments were carefully designed to reproduce as closely as possible the physiological features of the gastrointestinal tract emphasizing the flow and mixing processes found in the human small intestine.

Two configurations of the SIM were used: opened and closed

I. **Closed SIM configuration.** Chyme was static during the experiment. No net flow was induced by the peristaltic pump in the lumen side. The only movement of the chyme was imposed by the segmentation cuffs.

II. **Opened SIM configuration.** In the opened SIM configuration, both peristalsis and segmentation movements were induced in the lumen side, then chyme was recirculated several times during the experiment trough the inner side (membrane)
of the SIM to obtain representative data since the surface area of the small intestine is much larger than that of a tube with the same diameter.

Both experiments were done for at least one hour to obtain representative data on glucose absorption.

3.4.1. Absorption of glucose in the SIM

As food formulation plays a key point in the rate of absorption and delivery of molecules. A series of experiments were performed in the SIM to study (i) the effect of mixing and food formulation on glucose absorption and then (ii) the effect of the frequency of contractions and food formulation on absorptive processes through the membrane.

3.4.1.1 Effect of mixing and food formulation

The effect of mixing and food formulation, specifically viscosity, on glucose absorption rates was investigated in the closed SIM configuration (with no net flow in the lumen side). Experiments were performed using aqueous solutions, as a control, and the biopolymers: guar gum (0.1%, w/v) and CMC (0.1 and 0.5%, w/v) when segmentation and no segmentation take place in the SIM (Table 3.1). The biopolymer (0.1 – 0.5%, w/v) and glucose (55 mM) were homogeneously dispersed in distilled water using a magnetic stirrer for two hours to ensure the complete hydration of the biopolymer.

The biopolymer-glucose solution was placed in the dialysis membrane (approximately 400 mL) and in the outside tubing the active was collected. Distilled water (1.0 L) was used in the outer tube for all experiments. The experiment was run at room temperature by around 1 hour taking samples every 10 minutes. All samples were prepared the same day of use. Two
processing conditions were investigated to study the effect of mixing on the glucose absorption:

(i) **Mixing induced by segmentation contractions.** In this case local flow was induced by segmentation contractions (SM) by cuffs C1 and C2 (Figure 3.4a) alternatively with a 2 second inflation time, 2 second deflation time, 2 second delay and 0.50 barg inlet air pressure.

(ii) **No mixing or segmentation (stationary flow).** Neither local nor net flows were induced in the lumen side; no segmentation movements (Figure 3.4b). Absorption of glucose was done by facilitated diffusion to and through the interfacial membrane as a result of the concentration gradient.

The overall flow in the recipient side (outside of the membrane) was constant at 120 rpm for both experiments and was generated from a variable speed peristaltic pump (P2). Each experiment was carried out in triplicate.

**Table 3.1 Summary of the experiments designed to study the mixing effect**

<table>
<thead>
<tr>
<th>Biopolymer</th>
<th>Concentration (% w/v)</th>
<th>Mixing condition</th>
</tr>
</thead>
<tbody>
<tr>
<td>Control</td>
<td>0.0</td>
<td>Segmentation -2s</td>
</tr>
<tr>
<td></td>
<td></td>
<td>No segmentation</td>
</tr>
<tr>
<td>Guar Gum</td>
<td>0.1</td>
<td>Segmentation -2s</td>
</tr>
<tr>
<td></td>
<td></td>
<td>No segmentation</td>
</tr>
<tr>
<td>CMC</td>
<td>0.1</td>
<td>Segmentation -2s</td>
</tr>
<tr>
<td></td>
<td></td>
<td>No segmentation</td>
</tr>
<tr>
<td>CMC</td>
<td>0.5</td>
<td>Segmentation -2s</td>
</tr>
<tr>
<td></td>
<td></td>
<td>No segmentation</td>
</tr>
</tbody>
</table>

These experiments were done in the closed SIM configuration.
Chapter 3 - Materials and Methods

Figure 3.4 Photographs of the SIM (a) with and (b) without segmentation contraction cuffs

Intestinal Cell Set up Characteristics. The mass transfer cell (dialysis membrane) had a length of 0.5 m that could be extended and adjusted according to the experiment; the cuff length was 0.12 m, the inner tube and the outer tube had a diameter of 0.03 m and 0.05 m, respectively.

3.4.1.2 Effect of frequency of contractions and food formulation

As glucose absorption could be influenced by the wall movements of the small intestine (Macagno et al., 1982), the experiments described here were designed to study the effect of different mixing frequencies (1, 2 and 3s⁻¹) on glucose absorption.

To study the effect that the number of contractions per minute has on glucose absorption a series of experiments was carried out by changing the frequency at which segmentation contractions took place in the SIM.
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The formulation or viscosity of the luminal side was changed by using a range of biopolymers: guar gum, citrus pectin and CMC at different concentrations (Table 3.2) in a limited range of viscosities (0.1 - 3.0 Pa.s). The range of viscosities selected is in the range found by Marciani et al., (2000) from highly viscous meals after their transit through the stomach. The study was done when segmentation was applied to the opened SIM in which chyme was recirculated by means of a peristaltic pump (P1, Figure 3.2) during 1 hour through the inner side of the SIM to obtain representative data. Contrary to the previous experiment, here flow and mixing in the SIM were induced by peristalsis (net flow) and segmentation (local flow). The overall net flow in the lumen side was constant at $1.6 \times 10^{-4}$ m$^3$/s for all the experiments and was generated from a variable speed peristaltic pump. The segmentation movement responsible for mixing of the intestinal content was reproduced as described in section 3.3.3 by the occlusion of the tube with the use of inflatable bladders.

For this experiment, each biopolymer (guar, pectin and CMC) was dissolved in 2.0 L of glucose solution (55 mM) where they were homogeneously dispersed overnight by means of an overhead digital stirrer with a helical impeller at approximately 600 rpm.

The solution was placed in the dialysis membrane side (650 mL) and in the outside tubing the active - glucose - was collected. Distilled water (1.0 L) was used in the outer tube for all experiments. The experiment was run at room temperature by around one hour taking samples every 10 minutes.
Table 3.2 Summary of the experiments designed to study the effect of frequency of contractions

<table>
<thead>
<tr>
<th>Biopolymer</th>
<th>Concentration (% w/v)</th>
<th>Mixing frequencies</th>
</tr>
</thead>
<tbody>
<tr>
<td>Control</td>
<td>0.00</td>
<td>1s 2s 3s</td>
</tr>
<tr>
<td></td>
<td>0.25</td>
<td></td>
</tr>
<tr>
<td></td>
<td>0.50</td>
<td></td>
</tr>
<tr>
<td></td>
<td>0.65</td>
<td></td>
</tr>
<tr>
<td></td>
<td>0.75</td>
<td></td>
</tr>
<tr>
<td>Guar Gum</td>
<td>1.00</td>
<td>1s 2s 3s</td>
</tr>
<tr>
<td></td>
<td>2.00</td>
<td></td>
</tr>
<tr>
<td></td>
<td>2.50</td>
<td></td>
</tr>
<tr>
<td></td>
<td>3.00</td>
<td></td>
</tr>
<tr>
<td>Pectin</td>
<td>0.25</td>
<td>1s 2s 3s</td>
</tr>
<tr>
<td></td>
<td>0.50</td>
<td></td>
</tr>
<tr>
<td></td>
<td>1.00</td>
<td></td>
</tr>
<tr>
<td></td>
<td>1.25</td>
<td></td>
</tr>
<tr>
<td>CMC</td>
<td>0.25</td>
<td>1s 2s 3s</td>
</tr>
<tr>
<td></td>
<td>0.50</td>
<td></td>
</tr>
<tr>
<td></td>
<td>1.00</td>
<td></td>
</tr>
<tr>
<td></td>
<td>1.25</td>
<td></td>
</tr>
</tbody>
</table>

These experiments were done in the opened SIM configuration under segmentation conditions.

3.4.2 Density and viscosity measurements

The density of the viscous solutions was measured at least in triplicates by using a calibrated pycnometer at 25 °C. Distilled water was used as a reference.

The apparent viscosity of the biopolymers were determined using an AR1000 rheometer (TA instruments Newcastle, Delaware, USA) equipped with an acrylic plate geometry (0.06 m diameter). Each experiment was repeated in triplicate at 25 °C. Based on the literature survey, viscometry profiles were acquired at different shear rates ranging from 0.1 to 1000 (1/s) (Marciani et al., 2000). Zero-shear viscosity ($\eta_0$) values (in Pa.s or mPa.s) were calculated from the viscosity/shear-rate profiles for each biopolymer. The viscometry profiles for each concentration of guar, pectin and CMC as a function of the applied shear rate ($\dot{\gamma}$) are shown in Figure 3.5 (a-c).
Figure 3.5 Viscosity profiles of solutions of glucose (55 mM) added with (a) guar gum, (b) high methoxyl pectin and (c) CMC at various concentrations.

For aqueous solutions the viscosity and density were assumed to be the same as those for water at 25 °C: 8.904 x 10^{-4} kg/m³ CRC Handbook of Chemistry and Physics (1989) and 997.08 kg/m³ (Perry et al., 1973), respectively.

3.4.3 Glucose determination
Diffusion across the membrane was monitored at time intervals during one hour by measuring glucose concentration in the recipient side using the method of 3,5-dinitrosalicylic acid (DNS) reagent for reducing sugars (Miller, 1959). This method tests for the presence of
the free carbonyl group (C=O) in monosaccharides. The reaction involves the oxidation of the aldehyde functional group present in glucose to a carboxylate ion group under alkali conditions. Simultaneously, the DNS reagent is reduced to 3-amino-5-nitrosalicylic acid. For the determination of reducing sugars a UV-VIS spectrophotometer Libra S12 (Biochrom, Cambridge, UK) was used for measuring the absorbance of the standards and samples at 540 nm. All absorbance measurements were done at least in duplicates.

For glucose calibration curve (Figure 3.6) standards covering the range 0 - 5.0 mM in the final assay solution were used. The standards were assayed by the same procedure as the samples. Reducing sugar concentrations were expressed as glucose equivalents.

![Glucose calibration curve](image)

Figure 3.6 Glucose calibration curve in a range from 0.0 to 5.0 mM

### 3.4.4 Mass transfer coefficients

Overall mass transfer coefficients ($K$) were estimated by Equation 3.1.

$$N_l = K (c_i - c_\infty)$$

(3.1)

where $N_l$ is the molar flux (mmol/m$^2$ s) including both diffusion and convection and $c_i$ (mol/m$^3$) is the concentration at the interface of the same fluid as the bulk concentration $c_\infty$.
(mmol/m$^3$). The reciprocal of the overall mass transfer coefficient $K$ can be represented by the sum of the film resistances and that through the membrane and the recipient side (Equation 3.2)

$$\frac{1}{K} = \frac{1}{k_{bp}} + \frac{l_{mem}}{D_{mem}} + \frac{1}{k_{rep}}$$

(3.2)

where $k_{bp}$ is the biopolymer side mass transfer coefficient, $D_{mem}$ is the diffusion coefficient of nutrient through the membrane, $l_{mem}$ is the thickness of the membrane and $k_{rep}$ is the recipient side mass transfer coefficient.

To find the mass transfer enhancement from mixing it is important to calculate the Sherwood number (Equation 3.3)

$$Sh = \frac{k_{bp}L}{D_{AB}}$$

(3.3)

where $k_{bp}$ is the biopolymer side mass transfer coefficient, $L$ is the length scale (0.032 m) and $D_{AB}$ is the diffusion coefficient of glucose $6.9 \times 10^{-10}$ m$^2$/s (Perry et al., 1973).

### 3.4.5 Chemical Digestion in the SIM

Chemical digestion in the SIM involves the breakdown of macromolecules such as proteins, lipids and carbohydrates into small molecules, amino acids, fatty acids and sugars, respectively. Since glucose blood levels could be correlated in vitro to the starch digestion rates, this work was limited to starch-containing foods.
3.4.5.1 Starch preparation

Corn starch (0.5 – 3.0%, w/v) was previously gelatinized in a boiling water bath during 30 minutes with intermitted mixing. After gelatinization, the starch solution was cooled at room temperature.

3.4.5.2 Enzyme preparation

Pancreatic α-amylase (Type VI-B, ≥10 units/mg solid, A3176 Sigma) and 0.5 ml of amylglucosidase (from Aspergillus niger, ≥ 300 U/mL, aqueous solution, A7095, Sigma) were dissolved in a pancreatic mix solution (50 ml) at room temperature. The pancreatic mix solution was prepared by dissolving in 1 L of distilled water the following salts: NaCl (146mM), KCl (4.8mM), CaCl₂ (2.6mM), MgCl₂ (0.3 mM), ZnSO₄·7H₂O (4.1 μM). These concentrations are based on those reported by Lo Curto et al. (2011) and Madalari et al. (2010b) from the Institute of Food Research in Norwich, UK.

3.4.5.3 Gastric digestion

Gastric digestion was mimicked by reducing the pH of the corn starch solution from 5.5 to 2.5 with 1.0 M HCl. After 30 min, the solution was neutralized increasing the pH from 2.5 to 6.5 with 1.0 M NaHCO₃ or 0.1 N NaOH. BIS-Tris buffer (0.1 M, Sigma-Aldrich, UK) to adjust the pH of the samples at 6.5. For mimicking this step and considering that most of the starch hydrolysis occurs in the mouth and the intestine, no enzymes were added to the solution.

3.4.5.4 Intestinal digestion in the SIM

The starch was fed into the lumen side of the SIM. Starch digestion started when the pancreatic mix solution (50 ml) containing α-amylase (300 U/l) was pumped into the
membrane (~3 ml/min). The enzyme solution (25 or 50 ml) was freshly prepared for the digestion analysis by dissolving the pancreatic α-amylase (25 U/ml) and amyloglucosidase (3 U/ml) into the pancreatic solution for 10 minutes at room temperature. Starch digestion was done by using two in vitro models: (i) a stirred tank reactor and (ii) a small intestine model.

I. Stirred Tank Reactor (STR). Starch hydrolysis was carried out in a STR (Figure 3.7a) equipped with a Rushton turbine impeller and four baffles to create turbulence increasing the mixing. During the reaction, the impeller speed, temperature, and pH were kept constant at 300 ± 50 rpm, 25 ± 2 °C and 6.50 ± 0.05, respectively. The reactions were carried out in 250 ml of aqueous solutions containing 10 g/l corn starch.

II. Small Intestine Model (SIM). The digestion of starch was carried out in the inner tubing (membrane) or lumen side (Figure 3.7b) with 600 ml of liquid and semisolid food system containing 10 g/l corn starch. For these experiments, the closed-SIM configuration was used. The reaction was started by pumping the pancreatic mix solution into the reactor of the STR or in the lumen side of the SIM at ~3.0 ml/min (Figure 3.7 a,b). As before, two configurations were used for studying starch and food digestion: an open and a closed SIM.

Figure 3.7 In vitro models used for starch hydrolysis, (a) Stirred Tank Reactor (STR) and (b) Small Intestine Model (SIM).
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One of the innovations of the SIM compared with previous works is the implementation of an internal system to feed the enzyme into the SIM (similar to the pancreatic duct in the duodenum). As a first approach a long polypropylene tubing (3 mm of diameter) was introduced in the membrane section to feed the enzymes. This system allows changing the flow rate at which the enzyme is fed into the lumen side.

Food formulation was evaluated by adding guar (0.5%, w/v) to solutions of starch and white bread as model foods. For bread (50 g), the aqueous solution containing 0.5% (w/v) of guar was added into small pieces (~ 5 mm) simulating mechanical breakdown in mouth. In both cases, 1.0 L of the solution was prepared. Experiments were carried out in the closed-SIM configuration.

Preliminary experiments were done in the open-SIM configuration to study the effect of both flow rate and enzyme concentration on starch digestion. These experiments were carried out using corn starch solutions (1.0%, w/v) after gastric digestion section. For intestinal digestion, the enzyme solution (50 ml) was freshly prepared for the digestion analysis by dissolving the pancreatic α-amylase (2.5 and 25 U/ml) and amyloglucosidase (3 U/ml) into a pancreatic mix solution for 10 minutes at room temperature. The reaction was started by pumping the pancreatic solution into the SIM at 0.3 and 3.0 ml/min. The feeding flow rates at which the pancreatic solution was injected to the lumen side of the SIM were 0.3 and 3.0 ml/min. All combinations were tested to evaluate the effect of the way at which the enzymes were fed into the system.
Simulation of intestinal digestive processes is very challenging due to the difficulty to assess the small intestine environment and the variability of physiological conditions such as enzymes concentration and flow rates from person to person.

A number of conditions were studied to validate digestion and absorption processes in the SIM. As such, the flow of the enzyme solution together within the concentration of both enzyme and starch were studied in preliminary experiments as described in Table 3.3.

Table 3.3 Summary of the experiments done for studying starch hydrolysis

<table>
<thead>
<tr>
<th>Factors</th>
<th>Experiment 1</th>
<th>Experiment 2</th>
<th>Experiment 3</th>
</tr>
</thead>
<tbody>
<tr>
<td>Starch concentration</td>
<td>1.0% (w/v)</td>
<td>1.0% (w/v)</td>
<td>0.5% (w/v)</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>2.0% (w/v)</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>3.0% (w/v)</td>
</tr>
<tr>
<td>Enzyme concentration</td>
<td>10 U/ml</td>
<td>5 U/ml</td>
<td>15 U/ml</td>
</tr>
<tr>
<td></td>
<td>10 U/ml</td>
<td>10 U/ml</td>
<td></td>
</tr>
<tr>
<td></td>
<td>15 U/ml</td>
<td>15 U/ml</td>
<td></td>
</tr>
<tr>
<td>Feeding enzyme flow rate</td>
<td>1.5 ml/min</td>
<td>3.0 ml/min</td>
<td>3.0 ml/min</td>
</tr>
<tr>
<td></td>
<td>3.0 ml/min</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>4.5 ml/min</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Experiments done in the closed SIM configuration at pH 6.5 and room temperature at 2s segmentation. All experiments were performed in triplicate. No previous gastric digestion was done in these experiments.

Following the first section of glucose absorption, the effect of the food formulation and mixing frequencies were also evaluated (Table 3.4). Starch-containing foods were formulated by adding guar gum at different concentrations.
Table 3.4 Summary of the experiments done for studying the effect of food formulation on starch digestion

<table>
<thead>
<tr>
<th>Biopolymer</th>
<th>Concentration (%, w/v)</th>
<th>Mixing frequency</th>
</tr>
</thead>
<tbody>
<tr>
<td>Control</td>
<td>0.00</td>
<td>2s</td>
</tr>
<tr>
<td></td>
<td>0.25</td>
<td></td>
</tr>
<tr>
<td></td>
<td>0.50</td>
<td></td>
</tr>
<tr>
<td>Guar Gum</td>
<td>0.75</td>
<td>2s</td>
</tr>
</tbody>
</table>

Experiments done in the closed SIM configuration at pH 6.5 and room temperature. All experiments were performed in triplicate.

Glucose concentration was monitored in the lumen and the recipient side (outer tube). Samples (1.0 mL) were taken at time intervals to determine glucose absorption. The concentration of reducing sugars was calculated following the method of 3,5-dinitrosalicylic acid (DNS) reagent for reducing sugars measured samples absorbance using a UV-VIS spectrophotometer Libra S12 (Biochrom, Cambridge, UK) at 540 nm. For glucose calibration curve, standards covering the range 0 - 5mM in the final assay solution were used. The standards were assayed by the same procedure as the samples. Reducing sugar concentrations were expressed as glucose equivalents.

3.4.5.5 Data analysis

The starch hydrolysis results were analyzed according to the rate of digestion (lumen side) and absorption (recipient side) calculating the slope of the curve by linear regression using the method of least squares. These results were used later to calculate the overall mass transfer coefficients as described in section 3.4.4.
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**Kinetic parameters**

The *in vitro* kinetic parameters, based on Michaelis-Menten kinetics as described below for starch hydrolysis were estimated from the progress of the reaction. This relation assumes that the concentration of substrate (starch) is much bigger that the concentration of the enzyme.

The Michaelis-Menten equation in the differential form can be used to describe the dynamics of substrate depletion as described in Equation 3.4

\[
\frac{dS}{dt} = - \frac{V_{max} S}{K_m + S}
\]  
(3.4)

where \( S \) is the substrate concentration, and \( V_{max} \) and \( K_m \) are the maximal rate and Michaelis half saturation constant, respectively. Equation 3.4 can be readily integrated to obtain the integral form of the Michaelis-Menten (Equation 3.5.)

\[
K_m \ln \left( \frac{S_o}{S} \right) + S_o - S = V_{max} t
\]  
(3.5)

where \( S_o \) is the initial substrate concentration. The parameters of equation 3.5 were initially estimated by linearization and then recalculated using an explicit solution based on the *Lambert W* function (Goudar et al., 2004).

### 3.4.6. Flow visualization techniques

Particle imaging techniques measure the motion of a fluid on a small pre-determined region. Various techniques are available for flow visualisation in which markers are added at various densities. Two or more successive images are taken one after another and from the positions of the markers on the successive images; the flow field can be reconstituted.


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3.4.6.1 Planar Laser Induced Fluoresce (PLIF)

The PLIF technique is a non-intrusive approach to visualize the mixing mechanisms characterized by the tracer concentration - in this case Rhodamine 6G was selected for its fluorescence properties. The principle is based on the monotonic relationship between the tracer concentration and the fluorescence intensity (Hu et al., 2010).

Experimental cell configuration

The PLIF experiments were carried out in a Perspex rectangular cell (Figure 3.8), designed and implemented by Tharakan (2008) for this specific purpose. Within a length of 0.2 m and a height of 0.055 m, the cell containing an inner Perspex cylinder in the middle at which 20 cm of a transparent polypropylene tube was attached at both ending sides of the SIM (Figure 3.8 and 3.9).

![Experimental SIM set up including visualization cell for PLIF experiments.](image)

**Figure 3.8** Experimental SIM set up including visualization cell for PLIF experiments.
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PLIF/PIV system has a modular design comprised of a TSI Powerview system (TSI Inc. USA) which is made from the following: (i) a single 1000×1016 pixel, (ii) 8 bit Charged Coupled Device (CCD) camera (TSI PIVCAM 10-30, TSI Inc, USA); (iii) synchronizer, (iv) a dual head Nd-Yag laser emitting at 532 nm (New Wave Inc., USA); and (v) laser sheet optics. The synchronizer is controlled by a Dell Precision 620 workstation (Dell Corp. USA) that triggers the laser pulse and camera with the correct sequence and timing. PLIF images were captured by a high speed CCD camera within a very short time between them as a sequence.

Figure 3.9 PLIF Experimental Set up
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For these experiments, a -15 mm cylindrical lens and a 200 mm spherical lens were used, to give a minimum thickness of the laser sheet. The system was controlled by the INSIGHT 6.0 software.

After fixing the relative position of the SIM, the laser and the camera, three different concentrations of Rhodamine B previously dissolved in the correspondent liquid (water, CMC 0.5 or 1.0%, w/v) were filled in the stirred tank, then the fluorescence signals captured by the high speed camera were used for further calibration.

In the intestinal lumen, liquids are viscous with non-Newtonian characteristics. In these experiments, distilled water (control) and CMC solutions at two different concentrations (0.5 and 1.0%, w/v) were fed into the SIM to investigate the effect of the liquid viscosity on the mixing. The zero-shear viscosity of the solutions were 20 and 200 mPa.s for CMC 0.5% (w/v) and CMC 1.0% (w/v), respectively with a Reynolds number between 10 and 100. For the purposes of PLIF a single frame was captured each time and a sharp cut-off high-pass wavelength filter (ALP545, TSI Inc.) was used on the camera to eliminate the light from the laser and only allow fluorescent light from the tracer fluid to pass through. Images were captured at a rate of 2.0 Hz and the experiments were conducted for approximately 60 seconds.

Once the inner cylindrical system was homogenously filled with the distilled water and the CMC solution, the acrylic box was filled with tap water to reduce the light distortion due to the curved surface. 0.2 ml of fluorescent dye (Rhodamine 6G, Sigma Aldrich) was introduced as the tracer (at a concentration of 0.5 mg/l) into the fluid by a syringe at the centre of the
Perspex pipe before segmentation contractions started in the SIM. The rhodamine dye solution was made using CMC at the same concentration as the fluid used in the experiment. As indicated in Table 3.5, two mixing frequencies were evaluated for each solution: 1s\(^{-1}\) and 2s\(^{-1}\). The images were then processed with MATLAB and the Insight software.

Table 3.5 Summary of the experiments performed with the PLIP technique

<table>
<thead>
<tr>
<th>Biopolymer</th>
<th>Frequency of compression (segmentation)</th>
<th>1s</th>
<th>2s</th>
</tr>
</thead>
<tbody>
<tr>
<td>Control</td>
<td></td>
<td>Net flow and local flow</td>
<td>Net flow and local flow</td>
</tr>
<tr>
<td>CMC 0.5% (w/v)</td>
<td></td>
<td>Net flow and local flow</td>
<td>Net flow and local flow</td>
</tr>
<tr>
<td>CMC 0.5% (w/v)</td>
<td></td>
<td>Local flow</td>
<td>Local flow</td>
</tr>
<tr>
<td>CMC 1.0% (w/v)</td>
<td></td>
<td>Net flow and local flow</td>
<td>Net flow and local flow</td>
</tr>
<tr>
<td>CMC 1.0% (w/v)</td>
<td></td>
<td>Local flow</td>
<td>Local flow</td>
</tr>
</tbody>
</table>

3.4.6.2 Positron Emission Particle Tracking (PEPT)

PEPT is a non-invasive technique used to monitor motion inside opaque systems. PEPT is a unique technique that allows studying three dimensional flow phenomena as a function of both time and space. Due to its advantages, PEPT was used to study the flow dynamic in the SIM as a function of mixing and food formulation. These experiments were carried out at the Positron Image Centre at the University of Birmingham (UK).

Tracer properties. The radiotracer used in this research was adsorbed onto a 600 µm ion exchange resin particle and then coated with a layer of lacquer and paint to avoid the leaching of the radioactive isotopes into the fluid and to control the density to 1100 - 1200 kg m\(^{-3}\) (isokinetic with the fluid). The radiotracer used, \(^{18}\)F, has a sufficiently short half-life 109 min
that radiation does not persist in the system. The tracer was added to the fluid and used to track the fluid motion in the SIM.

**PEPT camera.** The motion of the tracer was tracked using an ADAC Forte positron camera (Parker et al., 2002) which employs two digital camera heads of size $500 \times 400 \text{ mm}^2$ operating in coincidence, the separation of the heads was adjusted to a maximum of 100 cm. The size of the camera field of view is therefore $800 \times 500 \times 400 \text{ mm}^3$. Each detector head contains a gamma digital camera comprising of a 16 mm thick NaI(Tl) scintillator, backed by an array of 55 photomultiplier tubes which are connected to analogue to digital converters and then to a computer.

**Experimental set up.** First, the lumen side of the SIM was placed between the detectors of PEPT dual camera leaving a gap of 0.55 m as shown in the Figure 3.10. Then the guar solution (0.5 or 1.0%, w/v) was fed into the membrane by means of a peristaltic pump (Watson-Marlow 323S/D). Once the system was filled with the solution the tracer previously dissolved in 5 ml of the same solution, it was injected with a syringe and a needle BD 16G1 (PrecisionGlide Needle No. 305197) in the tubing of the SIM approximately at 5 cm before the left exit. Experiments (Table 3.6) were performed on the SIM varying the fluid viscosity by addition of guar gum (0.5 - 1.0%, w/v). Factors such as mixing frequency and net flow on the mixing efficiency were studied. Each experiment was performed for at least 50 minutes in order to ensure sufficient sampling time. A summary of the experiments that were carried out is given in Table 3.6.
Table 3.6 Summary of the experiments performed with the PEPT camera

<table>
<thead>
<tr>
<th>Biopolymer</th>
<th>Frequency of compression (segmentation)</th>
<th>1s</th>
<th>2s</th>
</tr>
</thead>
<tbody>
<tr>
<td>Guar gum 0.5% (w/v)</td>
<td>Net flow and local flow</td>
<td>Net flow and local flow</td>
<td></td>
</tr>
<tr>
<td>Guar gum 0.5% (w/v)</td>
<td>Local flow</td>
<td>Local flow</td>
<td></td>
</tr>
<tr>
<td>Guar gum 1.0% (w/v)</td>
<td>Net flow and local flow</td>
<td>Net flow and local flow</td>
<td></td>
</tr>
<tr>
<td>Guar gum 1.0% (w/v)</td>
<td>Local flow</td>
<td>Local flow</td>
<td></td>
</tr>
</tbody>
</table>

From the PEPT experiments, 2-D and 3-D data of the position of the tracer versus time were obtained. The data obtained from the PEPT experiments were further processed using Matlab version 7.

3.4.7 Encapsulation of actives

To study stability of functional ingredients under gastrointestinal conditions and potential use in control delivery, riboflavin and alkaline phosphatase were encapsulated in gel particles and liposomes.
3.4.7.1 Gelled particles

Sodium alginate (1.0 - 3.0%, w/v), and riboflavin were homogeneously dispersed in distilled water by means of a magnetic stirrer. This solution was dropped using a syringe (0.7 mm of inner diameter) in a second solution, containing CaCl$_2$ (0.1, 0.3 and 0.5 mM) under gentle stirring. Microspheres (MS) were instantaneously produced by ionotropic gelation. The MS formed were allowed to stand in the solution for 15 minutes and then separated from the reaction mixture by filtration and washed with distilled water.

Entrapment efficiency

An indirect method was used to quantify the amount of active trapped in the MS (González-Rodríguez et al., 2002). Aliquots from the filtered solutions remaining after removal of the beads were assayed using a spectrophotometer (CECIL instrument CE 1020) at 665 nm. The amount of substance entrapped was calculated from the difference between the total amount of the colorant added and the amount found in the filtered solution.

Release of actives

In order to determine the amount of vitamin released from the MS of alginate, 1.0 g of each formulation was placed in a STR containing 20 ml of either gastric or intestinal fluid. The composition of each solution (gastric or intestinal) is described below (Rayment et al., 2009):

**Simulated Gastric Fluid (SGF):** NaCl 49.0 mM, KCl 11.6 mM, and CaCl$_2$ 3.6 mM, simulate conditions of the stomach and adjust pH to 2.0.

**Simulated Intestinal Fluid (SIF):** NaCl 111.0 mM, KCl 11.2 mM, CaCl$_2$ 2.0 mM and NaHCO$_3$ 16.5 mM, to mimic the pH of the small intestine, pH was adjusted to 7.0.

To simulate the release under gastric conditions, MS were placed in the SGF during 2 hours and then they were transferred to the SIF. After three hours in the SIF, the remaining MS
were placed in a buffer solution (pH 7.4). Samples (1.0 mL) were taken every hour and then analysed spectrophotometrically at 445 nm. The release was monitored during five hours. Percentage of cumulative release was calculated to obtain the dynamic profile release represented graphically. The release data were expressed as mean ± standard deviation based on three measurements. All experiments were done at room temperature.

3.4.7.2 Liposomes

Multilamellar vesicles (MLV) were prepared using the dry lipid film hydration method first developed by Bangham et al. (1965). Briefly, the neutral lipid Dipalmitoylphosphatidylcholine (DPPC) from Avanti polar lipids was dissolved in a 9:1 solvent mixture of chloroform and methanol. The solvent was evaporated on a rotary evaporator to obtain a dry film, which was hydrated with a solution of alkaline phosphatase (ALP) in 10mM Tris HCl buffer pH 7.4 (to give final lipid concentration 32 µmol/ml) in a round bottomed flask at 45 °C with intermittent vortexing. Unentrapped ALP was removed from the liposome suspension by repeated centrifugation decanting the supernatant and washing the pellet with 10mM Tris HCl until no ALP was detected in the supernatant.

Preparation of Gel loaded liposomes

Liposomes loaded with sodium alginate and the enzyme ALP where prepared using the thin lipid evaporation method described previously and hydrating with a mixture of 1% (w/v) sodium alginate containing 10mg/ml ALP. Following removal of unentrapped alginate/ALP the gel loaded liposomes were suspended in 200mM CaCl₂ for 30 min to induce gelation of the entrapped alginate. The diffusion of Ca²⁺ across the liposome bilayer was facilitated by increasing the temperature of the liposome suspension to the Tₘ of DPPC (42 °C) which, has
the effect of increasing bilayer permeability (Hong et al., 2008). The CaCl$_2$ was then removed by centrifugation and the liposomes were re-suspended in 10mM Tris HCl.

**Sizing and zeta potential measurements of liposomal dispersions**

The volume mean diameter (VMD) of the liposomes was measured using a Mastersizer (Malvern Instruments, Malvern, UK) at 25 °C, diluting 30µl of the dispersion to the appropriate volume with double distilled water. The zeta-potential, which is an indirect measurement of the liposome surface charge, was measured in 1.0mM Tris HCl pH 7.4 using a Zetamaster (Malvern Instruments, Malvern, UK).

**Entrapment efficiency**

The concentration of entrapped ALP was determined using the Bicinchoninic acid (BCA) assay (Sigma, Poole, UK). Quantification of protein in aqueous solutions using this assay was determined by the extent of reduction of Cu$^{2+}$ to Cu$^{+}$ which was measured colorimetrically using a UV-VIS spectrophotometer at 562 nm.

**Catalytic activity of ALP**

Catalytic activity of ALP was determined using an ALP activity kit (ABD Bioquest, California, USA). Briefly, p-nitrophenyl phosphate (pNPP) acted as the substrate which was hydrolysed by alkaline phosphatase releasing the chromomeric p-nitrophenol which was measured using a UV-VIS spectrophotometer at 450 nm. To quantify the protective effect of encapsulation, samples of ALP loaded liposomes, gel core liposomes and naked ALP were exposed to simulated gastric pH (pH 2) for up to 2 h. At selected time points samples were taken and the pH was raised to pH 8 by addition of 1.0M NaOH. The ALP was then released from the liposomes and gel core liposomes by treatment with 2% (w/v) sodium dodecyl
sulphate and 2% (w/v) sodium citrate. The catalytic activity of the released ALP was then measured as a function of time exposed to pH 2. The amount of ALP released while incubated at pH 2 was also monitored, samples were taken over a 2 h period, centrifuged and the ALP content of the supernatant was quantified using the BCA assay.

**pH Stability Study**

Samples of liposomes and gel core liposomes loaded with ALP were prepared and stored in 10mM Tris HCl maintained at pH 3.8, 7.4 and 10 at 4 °C. The particle size of the samples was measured (using the method described above) during storage at selected time intervals up to 10 days.

**Environmental Scanning Electron Microscopy (ESEM)**

The external morphology of the liposomes and the gel core liposomes was visualized using Environmental Scanning Electron Microscopy (ESEM). The internal morphology of the liposomes was analyzed by Cryogenic Scanning Electron Microscopy (Cryo-SEM). All ESEM analyses were performed with a Philips XL30 ESEM-FEG. The accelerating voltage was 3 kV and the working distance 5 mm. Sample preparation for the Cryo-SEM was performed as follows: Liposomes samples were frozen in liquid nitrogen slush (-196 °C) under vacuum. The samples were then fractured and coated with a thin layer of gold (approx. 10nm) to make the sample conductive. Finally, each sample was placed on a stage cooled to around -140 °C to be imaged.

**Statistical Analysis**

Comparison of means was conducted using a one-way analysis of variance (ANOVA) with post-hoc Tukey (HSD) test to identify significant differences (p<0.05) between data sets.
3.5 Conclusion

This chapter summarizes the experimental methods used throughout this work to fulfil the objectives proposed. A description of the chemicals and enzymes used was also included within this chapter. Glucose, starch and viscous polymers are some of the materials used to understand digestion and absorption processes occurring in the small intestine using an in vitro small intestine (SIM). A brief description of the SIM was also included with its main components. Basically, this chapter was divided accordingly to the following chapters including; (i) glucose absorption, (ii) starch digestion, (iii) flow visualization and (iv) encapsulation of actives.
CHAPTER 4 - ABSORPTION OF ACTIVES IN THE SIM

4.1. Introduction

The development of functional foods and novel drug formulations require a better fundamental understanding of digestion and absorption processes occurring in the GI tract. As the main portal of absorption of the GI tract is the small intestine, Chapters 4 and 5 are focused in understanding how mass transfer takes place in the SIM during absorption and digestion, respectively. Absorption of nutrients is needed not simply for energy but general metabolism and growth. Before absorption can take place in the gut, digestion of food is required to break polymers and other molecules into small fragments which can be absorbed through the epithelial cells of the small intestine.

Previous studies have shown that absorption of actives is influenced by the wall intestinal motility (Nguyen et al., 2008, Udaykumar et al., 2006, Macagno et al., 1982), however, little evidence exists to relate wall movements generated by peristaltic and segmentation contractions to mucosal transport. Since the small intestine is not subject to direct quantitative study (Macagno et al., 1982) due its location, studies on fluid motion and motility are quite challenging. As such, the development of in vitro models to study digestion and absorption is increasing. However, absorption models have structural and anatomical limitations that restrict their use in modelling these processes.

This chapter covers a systematic-quantitative study of the effect that mixing or segmentation contractions has on mass transfer of actives. As glucose is an essential marker in control dietary chronic disease such as diabetes, it was selected as the model molecule for experiments described in the following sections. The effect that mixing and formulation has on glucose absorption was also studied in the SIM. The chapter is divided into two sections in
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which the effect of: (i) mixing or segmentation and (ii) the frequency of segmentation contractions have on absorption were studied. In both sections, the impact of food formulation was also evaluated by adding viscous polymers such as guar, pectin and CMC to glucose-food models. The effect of mixing was evaluated in the closed-SIM configuration while the effect of frequency in the open-SIM configuration (see section 3.4).

4.2 Effect of mixing and food formulation on glucose absorption

4.2.1 Glucose absorption rates

To evaluate the influence of mixing on absorption a series of experiments applying segmentation contractions in the SIM were performed as described in section 3.4.1.1. The experiments were carried out at room temperature for one hour in the SIM configuration, in which both ends of the lumen side (membrane) were closed, i.e. no recirculation of the chyme was done. Segmentation contractions pattern used in this section was as follows: 2s cuff inflation, 2s cuff deflation and 2s delay for each cuff.

The increase in glucose over time on the recipient side (outside of the membrane) has been plotted in Figure 4.1 (a) and (b) respectively, for aqueous glucose solutions (control) and glucose-guar (0.1%, w/v) solutions with low zero shear rate viscosity (1 - 2 mPa.s). Results showed that glucose absorption was accelerated when segmentation occurred in the SIM. After one hour, glucose concentration in control solution (without guar) was:

- Under segmentation conditions, almost 6.0 mM, around 10% of the initial glucose concentration (55 mM) as shown in Figure 4.1 (a) and
- Without segmentation, only 4.0 mM that represents only 7.0% of 55 mM (initial concentration) as Figure 4.1 (a) shown.
Chapter 4. Absorption of actives

Furthermore, this tendency was observed for guar (0.1%, w/v) when the zero shear viscosity of the solution was increased up to 2.0 mPas (Figure 4.1b). In both cases (control and guar) segmentation increased the absorption of glucose, conversely as the viscosity of the solution increased the difference between segmentation and no segmentation was clearly reduced. For example, the increase of absorption for aqueous glucose solutions under segmentation was up to 30%, whereas for guar-containing solutions, absorption was only improved around 10%.

Figure 4.1 Absorption of glucose in (a) aqueous solutions (1.0 mPa.s) and (b) guar gum solutions (2.0 mPa.s) when segmentation and no segmentation motion take place in the SIM.
The effect of CMC (0.1 and 0.5%, w/v) on glucose absorption process is shown in Figure 4.2. There was a clear reduction on glucose absorption when the viscosity of the solution increased from 20 to 200 mPas. After one hour, glucose concentration in 0.1% (w/v) of highly viscosity CMC solution was:

- Under segmentation conditions, almost 4.5 mM, around 8.0% of the initial concentration of glucose in the lumen (55 mM) and
- With no segmentation, only 3.55 mM; 6.5% of the total glucose in the lumen side of the SIM.

Nevertheless, these figures were significantly reduced when 0.5% (w/v) of highly viscosity CMC was added to the glucose solution:

- Under segmentation after one hour, glucose concentration in the recipient side was 1.5 mM, about 3.0% of the concentration of glucose in the lumen (55 mM) and
- With no segmentation, glucose reached only 0.3 mM, less than 1.0% of the glucose present in the lumen side of the SIM at the beginning of the experiment.

In both cases, segmentation increased absorption of glucose; the difference between segmentation and no segmentation was more significant as the viscosity of the solution increased.
Figure 4.2 Absorption of glucose in 0.1 and 0.5% (w/v) CMC solutions with zero shear viscosities of 20 mPa.s and 200 mPa.s, respectively, when segmentation and no segmentation motion take place in the SIM.

Relative to the control (water) under segmentation, glucose absorption was reduced up to 20 and 75% when 0.1 and 0.5% (w/v) of CMC were added to the solution of glucose, respectively. Furthermore, when no segmentation (diffusive transfer) was applied to the solution containing 0.5% (w/v) of CMC, the biopolymer restricts mass transport and the concentration of glucose was close to zero (Figure 4.2).

Overall, glucose was rapidly absorbed when segmentation contractions occur (convective transport by mixing), however, passive diffusion following Fick’s law took place when no segmentation contractions were applied. The greater the viscosity of the solution is, the lower the absorption. This is thought to be caused by decreased mixing in the lumen content.

Segmentation has a significant effect on glucose absorption rate increasing up to 30% for aqueous solutions. On the contrary, when viscous solutions were tested the impact of
segmentation was impaired and the difference between applying segmentation (convective transport) or not applying segmentation (diffusion transport) was only 10 - 20% for guar and CMC solutions. Since addition of CMC produced solutions with higher viscosity than guar solutions, CMC was more effective for controlling and delaying glucose absorption in the SIM. To compare the effect that each polymer has on glucose absorption experiments with similar viscosities must be done.

The glucose absorption rates (µM/s) were estimated from the slope of the curves in Figures 4.1 and 4.2 and are given in Table 4.1. Only the linear region of the curve was used to calculate the slope for each data series, i.e. from \( t_0 = 10 \) to \( t = 60 \) min, once the flow was fully developed. The concentration at time zero was not included as the actual position of zero is unknown. Triplicates were obtained for each condition.

The results indicate that both segmentation motion and viscosity have a significant effect on glucose absorption rates (Table 4.1). Whereas segmentation motion seems to increase the rates of glucose absorption, the increase in viscosity (from 1 to 200 mPa.s) resulting in a significant decrease on glucose concentration in the recipient side. The latter effect was even more drastic when no segmentation was applied to the SIM. As concentration of guar was increased and therefore the viscosity increased the effect of segmentation was reduced.

Table 4.1 Glucose absorption rates for experimental solutions.

<table>
<thead>
<tr>
<th>Biopolymer</th>
<th>Concentration (%)</th>
<th>Zero shear viscosity ± std (mPa.s)</th>
<th>Absorption rate ± std (µM/s)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>Segmentation</td>
<td>No segmentation</td>
</tr>
<tr>
<td>Control</td>
<td>0.0</td>
<td>1.0 ± 0.2</td>
<td>1.41 ± 0.19</td>
</tr>
<tr>
<td>Guar gum</td>
<td>0.1</td>
<td>2.0 ± 0.4</td>
<td>1.40 ± 1.02</td>
</tr>
<tr>
<td>CMC</td>
<td>0.1</td>
<td>20.0 ± 0.2</td>
<td>1.20 ± 0.14</td>
</tr>
<tr>
<td>CMC</td>
<td>0.5</td>
<td>200.0 ± 0.1</td>
<td>0.02 ± 0.26</td>
</tr>
</tbody>
</table>
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Experiments were performed in the closed-SIM configuration.

These results are in agreement with previous studies done in vitro by Tharakan et al., (2010) and in vivo by Blackburn et al., (1984), in which the addition of guar gum to glucose solutions fed to humans reduced the increase in blood glucose level, resulting in only 50% of that observed without guar. Blackburn et al. (1984) proposed that the mechanism of reduction is by inhibiting the effects of intestinal motility on fluid convection. Possible mechanisms are that viscous polysaccharides act by delaying gastric emptying and thus slow delivery to the small intestine and reduce the rate at which glucose is absorbed from the intestine (probably due to a reduced mixing efficiency).

The data obtained here can be compared with absorption rates in the human small intestine from Blackburn, et al. (1984). The rates of glucose absorption were estimated by the slope of the plasma glucose curves developed in the first 30 min period after ingestion of 50 g glucose in 250 ml of an orange drink with and without 9 g of guar gum (Figure 2.13). The absorption rates were:

- 1.85 \( \mu \text{M/s} \) for drinks without guar gum and
- 0.90 \( \mu \text{M/s} \) when guar was added to the solution of glucose (6.9 mM)

A significant (p<0.05) reduction (50%) of the rate of absorption in blood glucose levels was observed in humans after ingestion of drinks supplemented with 9 g of guar (36 g/l).

Although the concentration of glucose and guar were higher than those used in this research, a ‘good’ correlation between experiments carried out in the human small intestine and in the SIM can be established. The two sets have the same order of magnitude. However, it is important to remember this is a roughly comparison that could serve as a reference to correlate experimental data from the SIM to those obtained in vivo, in which glucose absorption is a carried-mediated transport that follows an electrochemical concentration
gradient in the cellular membrane of the epithelial cells. The intestinal epithelial cell membrane is a complex structure that consists of a double layer of lipid molecules containing embedded proteins responsible for transporting glucose from the cell to the blood vessels. However, the SIM used in this work is comprised by a semi-permeable membrane that simplifies the anatomical features of the intestinal wall.

### 4.2.2 Overall mass transfer coefficients (OMTC)

The overall mass transfer rate is given by equation 4.1, assuming that the lumen is a well-mixed solution and the mass transferred is proportional to the concentration difference and the interfacial area (Cussler, 2009).

\[
N_1 = K(c_{lt} - c_l)
\]

(4.1)

where \(N_1\) is the flux (mmol/m\(^2\)s), \(K\) is the mass transfer coefficient (m/s) and \(c_{lt}\) and \(c_l\) (mol/m\(^3\)) are the concentrations at the interface and in the bulk solution, respectively.

Considering for example, the case of the aqueous glucose solution under segmentation, the rate of glucose transferred is equal to \(1.41 \times 10^{-3}\) mM/s, from the slope of the curve in Figure 4.1(a), the membrane surface area was 0.05 m\(^2\) with a diameter of 0.03 m and the recipient side total volume was 0.001 m\(^3\). By dividing the rate of glucose transferred by the membrane surface area, the molar flux of this system is found to be \(2.94 \times 10^{-2}\) mmol/m\(^2\)s.

The overall mass transfer coefficient (\(K\)) can be estimated from equation 4.1 with a concentration difference of 55 mM, i.e. \(K\) is equal to \(5.35 \pm 0.07 \times 10^{-7}\) m/s (water under segmentation) and \(3.09 \pm 0.12 \times 10^{-7}\) m/s (water without segmentation). Similarly, \(K\) was estimated for guar and CMC.

The overall mass transfer coefficients (OMTC) for glucose absorption are presented in Table 4.2. The maximum OMTC were found for water (control) under segmentation conditions.
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This was expected as in this low viscosity solution, the resistance to mass transfer from the bulk of the fluid to the surface of the membrane was minimum. The OMTC decreased approximately 15% when the fluid viscosity increased ten-fold. On the other hand, when viscosity increased from 1.0 mPa.s (water) to 200 mPa.s (CMC 0.5%, w/v), K was reduced up to 90%.

Table 4.2 Overall Mass Transfer Coefficients (OMTC) for glucose absorption.

<table>
<thead>
<tr>
<th>Biopolymer</th>
<th>Concentration (%)</th>
<th>Zero shear viscosity ± std (mPa.s)</th>
<th>OMTC ± std (10⁻⁷ m/s)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>Segmentation</td>
<td>No segmentation</td>
</tr>
<tr>
<td>Control</td>
<td>0.0</td>
<td>1.0 ± 0.2</td>
<td>5.35 ± 0.07</td>
</tr>
<tr>
<td>Guar gum</td>
<td>0.1</td>
<td>2.0 ± 0.4</td>
<td>5.31 ± 0.39</td>
</tr>
<tr>
<td>CMC</td>
<td>0.1</td>
<td>20.0 ± 0.2</td>
<td>4.55 ± 0.05</td>
</tr>
<tr>
<td>CMC</td>
<td>0.5</td>
<td>200.0 ± 0.1</td>
<td>0.76 ± 0.06</td>
</tr>
</tbody>
</table>

Again these values can be compared with the limited in vivo data from the rates of glucose absorption calculated in section 4.2.1 for the human small intestine data (Blackburn et al., 1984). The OMTC was estimated as 5.47 x10⁻⁷ m/s for solutions without guar and 2.91 x 10⁻⁷ m/s, when 9 g of guar were added. The length of the small intestine was assumed to be 6.0 m with area of 0.575 m² and a volume of 0.012 m³. This assumes that everything is fully filled and the area of the small intestine is equal to a pipe with an average diameter of 0.03 m.

Figure 4.3 shows the effect of viscosity on mass transfer of glucose under both segmentation and no segmentation conditions. As it is shown in Figure 4.3, when viscosity increases from 0.001 to 0.200 Pa.s the mass transfer coefficient falls drastically from 5.35 x 10⁻⁷ m/s to nearly zero.
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4.2.3 Mass transfer coefficients in the biopolymer side

As Figure 4.4 shows, $K$ consists of three resistances: the biopolymer side $k_{bp}$ (donor), the membrane and the water film (recipient side).

The local mass transfer coefficient ($k_{bp}$) in the lumen or biopolymer side was calculated based on Equations 4.2 - 4.4 (Tharakan, 2008). This relation is based on the resistance that the membrane and the recipient side water film contribute to the OMTC ($K$). The last two terms in equation 4.3 can be considered as the resistance from the system $R_{system}$ (membrane and
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recipient side film). For these experiments the \( R_{\text{system}} \) was found to be \( 1.86 \times 10^6 \) s/m water under segmentation. This value was calculated as the inverse of the mass transfer coefficient found for water (\( 5.35 \times 10^{-7} \) m/s, Table 4.2). For fluids containing guar gum or CMC, \( k_{bp} \) was estimated by Equation 4.4.

\[
\frac{1}{k} = \frac{1}{k_{bp}} + \frac{l_{\text{mem}}}{D_{\text{mem}}} + \frac{1}{k_{repl}} \quad (4.2)
\]

\[
R_{\text{system}} = \frac{l_{\text{mem}}}{D_{\text{mem}}} + \frac{1}{k_{repl}} \quad (4.3)
\]

\[
k_{bp} = \frac{1}{\left(\frac{k}{k_{repl}}\right) - R_{\text{system}}} \quad (4.4)
\]

where \( k_{bp} \) is the biopolymer side mass transfer coefficient, \( D_{\text{mem}} \) is the diffusion coefficient of nutrient through the membrane, \( l_{\text{mem}} \) is the thickness of the membrane and \( k_{repl} \) is the recipient side mass transfer coefficient.

The range of mass transfer coefficients in the lumen side (biopolymer side) are shown in Table 4.3 under segmentation conditions.

Table 4.3 Local mass transfer coefficients of the biopolymer side from different viscous materials.

<table>
<thead>
<tr>
<th>Biopolymer (w/v)</th>
<th>( k_{bp} ) (( 10^{-6} ) m/s)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Guar_0.1%</td>
<td>3.22 ± 1.32</td>
</tr>
<tr>
<td>CMC_0.1%</td>
<td>1.49 ± 0.10</td>
</tr>
<tr>
<td>CMC_0.5%</td>
<td>0.01 ± 0.02</td>
</tr>
</tbody>
</table>

These results demonstrate that transport of actives to the membrane can be controlled by the properties of the chyme under physiological conditions. As concentration of CMC is increased, under segmentation conditions, mixing and flow at the membrane surface is decreased resulting in a reduction of \( k_{bp} \) from 1.5 to \( 0.01 \times 10^{-6} \) m/s.
To quantify the mass transfer enhancement of the system with convection compared to diffusion the Sherwood number \((Sh)\) was estimated by Equation 4.5.

\[
Sh = \frac{k_{bp} L}{D_{AB}} \tag{4.5}
\]

where \(k_{bp}\) is the biopolymer side mass transfer coefficient (Table 4.3), \(L\) is the length scale (0.032 m) and \(D_{AB}\) is the diffusion coefficient of glucose \(6.9 \times 10^{-10} \text{ m}^2/\text{s}\).

In addition, the average Reynolds number \((Re)\) of the fluids in the lumen or biopolymer side was calculated by Equation 4.6.

\[
Re = \frac{\rho v D}{\mu} \tag{4.6}
\]

where \(\rho\) is the density of the solution, \(v\) is the velocity of the fluid \((v = 0.06 \text{ m/s})\), \(D\) is the diameter of the membrane (0.03 m) and \(\mu\) is the viscosity of the solution (Table 4.1).

To calculate the \(Re\) of each solution, the maximum velocity (0.06 m/s) was estimated using the volume displaced by segmentation contractions at 2s. The velocity was calculated by assuming that the volume displaced by the cuff during each contraction is equal to \(9.65 \times 10^{-5} \text{ m}^3\). This value was estimated from the volume of a cylinder with a diameter of 0.03 m (membrane) and length of 0.12 m (length of the cuff in the SIM) as shown in Figure 4.5. By dividing the volume by the time each contraction is applied (2s), the volumetric flow was obtained \((4.83 \times 10^{-5} \text{ m}^3/\text{s})\). Finally, the mean velocity was estimated by dividing this volumetric flow by the cross sectional area of the flow \((7.07 \times 10^{-4} \text{ m}^2)\).
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Figure 4.5 A diagram of a segment of the SIM that represents the volume displace forwards by an inflated cuff (0.12 m length) when segmentation occurs.

The Sherwood numbers ($Sh$) against the Reynolds numbers ($Re$) for experiments when segmentation took place are shown in Figure 4.6. As the Reynolds number increases the Sherwood number increases as well, indicating that for low viscosity fluids mass transfer phenomena take place by convective forces. Mixing brings material (glucose) to the wall of the lumen side reducing the unstirred layer and resistance of the membrane to the transport of actives molecules. In contrast, diffusion occurs more likely when the effect of mixing is impaired by the viscosity of the fluid in the lumen side increasing the unstirred layer in the biopolymer side.

A suitable correlation (Treybal, 1980) was also used to estimate the $Sh$ inside circular pipes with similar dimensions and fluid flow than those studied in the SIM. Results are summarized and compared in Table 4.4 and Figure 4.6. Interestingly, experimental and theoretical values of $Sh$ were similar.
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Table 4.4 Experimental and theoretical Sherwood numbers estimated for the SIM and a circular pipe with similar fluid motion.

<table>
<thead>
<tr>
<th>Biopolymer</th>
<th>(\text{Re}_n)</th>
<th>(*\text{Sc})</th>
<th>(\text{Sh}_{\text{exp}})</th>
<th>(*\text{Sh}_{\text{the}})</th>
</tr>
</thead>
<tbody>
<tr>
<td>Control</td>
<td>2050</td>
<td>1300</td>
<td>650</td>
<td>140</td>
</tr>
<tr>
<td>Guar (0.1%, w/v)</td>
<td>900</td>
<td>3000</td>
<td>145</td>
<td>94</td>
</tr>
<tr>
<td>CMC (0.1%, w/v)</td>
<td>90</td>
<td>30000</td>
<td>66</td>
<td>30</td>
</tr>
<tr>
<td>CMC (0.5%, w/v)</td>
<td>90</td>
<td>300000</td>
<td>4</td>
<td>9</td>
</tr>
</tbody>
</table>

\(\text{Sc} = \frac{\mu}{\rho D}\) where \(\mu\) is the viscosity, \(\rho\) is the density and \(D\) is the diffusivity of glucose

\(*\text{Sh} = 0.023 \text{Re}^{0.83} \text{Sc}^{1/3}\) (Treybal, 1980).

Figure 4.6 Experimental and theoretical values of the Sherwood numbers for water, guar (0.1%, w/v) and CMC (0.1 and 0.5%, w/v).

To estimate conditions more like those found in the small intestine where oscillatory flow is produced by wall movements, a suitable correlation between oscillatory Reynolds number \(\text{Re}_o\) and the net flow Reynolds number \(\text{Re}_n\) was done. Oscillatory Reynolds numbers were calculated by Equation 4.7 developed according to Harvey et al., (2003).
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\[ Re_o = \frac{\rho 2\pi f x_o}{\mu} \]  

(4.7)

where \( \rho \) is the fluid density, \( f \) the frequency of oscillation (2s), \( x_o \) is the centre-to-peak amplitude (0.12 m) and \( \mu \) the viscosity of the fluid.

Results are summarized and compared in Table 4.5. Reynolds and Sherwood numbers were increased by ten-fold, when the oscillatory flow was used.

Table 4.5 Reynolds and Sherwood numbers calculated from the net (\( Ren \)) and oscillatory flows (\( Reo \)).

<table>
<thead>
<tr>
<th>Biopolymer</th>
<th>( Re_n )</th>
<th>( Sh_n )</th>
<th>( Re_o )</th>
<th>( Sh_o )</th>
</tr>
</thead>
<tbody>
<tr>
<td>Control</td>
<td>2050</td>
<td>140</td>
<td>25000</td>
<td>1133</td>
</tr>
<tr>
<td>Guar (0.1%, w/v)</td>
<td>900</td>
<td>94</td>
<td>11300</td>
<td>760</td>
</tr>
<tr>
<td>CMC (0.1%, w/v)</td>
<td>90</td>
<td>30</td>
<td>1133</td>
<td>240</td>
</tr>
<tr>
<td>CMC (0.5%, w/v)</td>
<td>90</td>
<td>9</td>
<td>113</td>
<td>77</td>
</tr>
</tbody>
</table>

Fluid motion in the small intestine is determined by the intestinal contractions generated during peristalsis and segmentation. Segmentation contractions are local constrictor waves that induce powerful radial mixing of the lumen contents enhancing mass transfer across the epithelium (Macagno and Christensen, 1980).

In conclusion, the results presented in this section suggest that segmentation enhanced mass transfer processes in the SIM. Nevertheless, mixing was impaired when the viscosity of the solution was increased as a result of a significant reduction in the rates of glucose absorption and the OMTC was achieved for viscous test food models.
4.3. Effect of mixing frequency and food formulation

Having established the importance of flow and mixing on mass transfer, the following experiments were designed to study the effect that frequency of contractions have on mass transfer during glucose absorption in the SIM. One of the advantages of this *in vitro* system is that different patterns or sequence of contractions in a physiological context can be studied by changing inflation time and delay between contractions (section 3.3.3). As a result, segmentation contractions can vary in intensity, duration and number of contractions per minute. The results presented in this section were obtained by using three types of segmentation contractions lasting from 1 to 3 seconds. The duration and the number of contractions per minute were changed accordingly to the patterns showed on Figure 4.8, resulting in three different frequencies:

1. 1s delay, 1s inflation, 1s deflation 20 contractions/minute (0.333 s\(^{-1}\))
2. 2s delay, 2s inflation, 2s deflation 10 contractions/minute (0.166 s\(^{-1}\))
3. 3s delay, 3s inflation, 3s deflation 7 contractions/minute (0.116 s\(^{-1}\))

The inflation and deflation terms refer to the time at which each inflatable cuff was supplied with compressed air (inflation) or vacuum (deflation) to generate segmental movements in the lumen side. The range of frequencies selected for these experiments were set in a physiological scenario in which 10-12 contractions per minute take place in the duodenum, decreasing in the jejunum and ileum until 7-8 in normal subjects (Guyton and Hall, 2006, Levy et al., 2006). An extreme case of study (20 contractions per minute) was selected to mimic gastrointestinal disorders such as irritable bowel syndrome, in which contractions or movements could be twice than those found in normal conditions.
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Figure 4.7  Deformation profiles of segmentation contractions applied to the SIM, in black the movements of cuff 1 and in blue the contractions of cuff 2, represented as ‘1’ and ‘2’, respectively, for 1 - 3 seconds. Each segmentation contraction has a symmetric delay, inflation and deflation times.

These values resulted in three different segmentation contractions profiles varying in intensity and duration. For 1s, the mixing or segmentation contraction was shallow; the inflation time was short and the cuff was not fully inflated applying a weak compression on the tube section of the SIM as demonstrated later in Chapter 6 section 6.2.2. Since this type of contraction was produced every second, the number of contractions per minute was approximately 20. In contrast, when cuffs were inflated for about 2s the contraction was deeper than the contractions generated every second. When segmentation contractions were applied every 2s, 10 pulses per minute were produced. The last type of contraction was the deepest with 3s of inflation, however, because the delay between contractions was extended to three seconds, around 7 pulses per minute were generated. During these experiments, recirculation of the lumen contents was done by using a peristaltic pump (open SIM
configuration). The net flow rate in the lumen side was $1.2 \times 10^{-5} \text{ m}^3/\text{s}$ (12 ml/s) resulting in an average velocity of $1.6 \times 10^{-4} \text{ m/s}$.

The effect of food formulation was evaluated by using three viscous polysaccharides: guar gum, pectin and carboxymethyl cellulose (CMC). The viscosity of the solution was modified by increasing the concentration of the biopolymer added to a solution of glucose used as a model food within an initial concentration of 10 g/L (~55 mM). Experiments were designed to have four solutions of each polymer with similar range of zero shear viscosities. The concentration of guar, pectin and CMC was adjusted to have solutions of low (0.01 - 0.1 Pa.s), medium (0.2 - 0.5 Pa.s) and high (>1.0 Pa.s) viscosities. The final range of viscosities was from 0.01 to 3.0 Pa.s. This range was selected according to the zero-shear viscosities of the chyme measured \textit{in vivo} by Marciani et al. (2000). These authors found that the viscosity of model meals added with locust bean gum decreased from 11 to 2 Pa.s after ingestion, due to the salivary and gastric secretions. Accordingly, the chyme enters to the duodenum within the range of viscosities chosen for the following experiments. All experiments were performed in duplicate and in a randomized order.

4.3.1 Glucose absorption rates: effect of frequency of contractions and formulation

4.3.1.1 Glucose absorption rates for guar gum solutions

The first biopolymer studied to modify the formulation of the test food was guar gum for a range of concentrations between 0.25 and 0.75% (w/v), densities close to water and zero-shear viscosities from 0.02 to 3.2 Pa.s (Table 4.6). Among the polymers used in this work, guar had the wider range of viscosities with a small change in concentration. As the concentration of guar increased the zero-shear viscosity increased as well, small doses of guar
added to the glucose liquid meal had great influence on the zero-shear viscosity of the solution.

Table 4.6 Physical properties of the experimental guar gum-glucose solutions.

<table>
<thead>
<tr>
<th>Assay</th>
<th>Concentration of guar (g/L)</th>
<th>Density (kg/m$^3$)</th>
<th>Zero-shear viscosity (Pa.s)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>2.50</td>
<td>1002.00 ± 0.33</td>
<td>0.0222 ± 0.0018</td>
</tr>
<tr>
<td>2</td>
<td>5.00</td>
<td>1003.18 ± 0.43</td>
<td>0.4108 ± 0.0296</td>
</tr>
<tr>
<td>3</td>
<td>6.25</td>
<td>1004.25 ± 0.44</td>
<td>1.2090 ± 0.0961</td>
</tr>
<tr>
<td>4</td>
<td>7.50</td>
<td>1006.01 ± 0.04</td>
<td>3.192 ± 0.1982</td>
</tr>
</tbody>
</table>

The guar-containing glucose solution (650 mL) was fed to the lumen or membrane side of the SIM. Absorption was monitored in the recipient side containing 1.0 L of distilled water every ten minutes during one hour. Segmentation contractions were produced accordingly to the patterns shown in Figure 4.7 every 1, 2, or 3 seconds. The results for glucose absorption in solutions added with guar gum are shown in Figure 4.8.

An evident reduction on absorption was observed as the viscosity of the solution increased. Surprisingly, the rates of glucose absorption were very similar and no significant difference was observed among frequencies tested for all range of concentrations of guar. Overall, glucose absorption decreased by increasing the concentration of guar added to the test food as a result of the increase in viscosity.

Since no significant differences were observed for guar, in dilute solutions of pectin (1.0%, w/v) and CMC (0.5%, w/v) only two frequencies were used (1s and 2s).
Figure 4.8 Glucose absorption in the SIM for solutions supplemented with guar gum at various concentrations: (a) 0.25, (b) 0.50, (c) 0.65 and (d) 0.75% (w/v) in a glucose solution (1.0 w/v) under segmentation contractions produced every 1, 2 or 3 seconds.

4.3.1.2 Glucose absorption rates for citrus pectin solutions

Pectin was also used to modify the formulation of the digesta. In this case, however, the range of concentrations was threefold higher than guar from 10 to 30 grams per litre of digesta (glucose solution) to reach the same range of viscosities established, like guar densities were close to water (Table 4.7).
Table 4.7 Physical properties of glucose-pectin solutions.

<table>
<thead>
<tr>
<th>Assay</th>
<th>Concentration of pectin (g/L)</th>
<th>Density (kg/m³)</th>
<th>Zero-shear viscosity (Pa.s)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>10</td>
<td>1004.79 ± 0.04</td>
<td>0.0498 ± 0.0217</td>
</tr>
<tr>
<td>2</td>
<td>20</td>
<td>1008.91 ± 0.24</td>
<td>0.2530 ± 0.0770</td>
</tr>
<tr>
<td>3</td>
<td>25</td>
<td>1010.63 ± 0.07</td>
<td>0.7133 ± 0.0607</td>
</tr>
<tr>
<td>4</td>
<td>30</td>
<td>1012.59 ± 0.15</td>
<td>1.9265 ± 0.1039</td>
</tr>
</tbody>
</table>

Absorption of glucose solutions containing pectin at various concentrations was monitored in the recipient side over time (60 minutes) at 1s, 2s, and 3s as shown in Figure 4.9.

**Figure 4.9** Glucose absorption in the SIM for solutions added with high methoxyl pectin at various concentrations: (a) 1.0, (b) 2.0, (c) 2.5 and (d) 3.0% in a glucose solution (1.0%, w/v) under segmentation contractions at 1, 2 and 3 seconds.
Similar to guar, the results showed that for food structured with pectin at various concentrations (1.0 - 3.0%, w/v), the number of contractions per minute and the intensity of the contraction gave rise to similar glucose absorption levels.

### 4.3.1.3 Glucose absorption rates for CMC solutions

The last biopolymer tested to study the effect of the mixing frequencies was CMC. As with guar and pectin, viscosity of the glucose solution increased as the concentration of CMC increased. In this case the range of concentrations was closer than those used with guar. The concentration of CMC that fixes the zero-shear viscosities previously selected was in the range between 0.25 to 1.25% (w/v), with densities close to water (Table 4.8).

<table>
<thead>
<tr>
<th>Assay</th>
<th>Concentration (%, w/v)</th>
<th>Density (kg/m³)</th>
<th>Zero-shear viscosity (Pa.s)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>0.25</td>
<td>1002.06 ± 0.03</td>
<td>0.0854 ± 0.0025</td>
</tr>
<tr>
<td>2</td>
<td>0.50</td>
<td>1002.88 ± 0.07</td>
<td>0.2698 ± 0.0004</td>
</tr>
<tr>
<td>3</td>
<td>1.00</td>
<td>1004.92 ± 0.14</td>
<td>0.5314 ± 0.0075</td>
</tr>
<tr>
<td>4</td>
<td>1.25</td>
<td>1006.01 ± 0.05</td>
<td>1.4503 ± 0.0410</td>
</tr>
</tbody>
</table>

Interestingly, when CMC was used the absorption of glucose seems to be affected by the frequency of segmentation contractions as shown in Figure 4.10. At 1s, the absorption of glucose was dramatically reduced compared with the other two frequencies, 2 and 3s, for concentrations equals or higher than 10 g of CMC per litre of digesta (Figure 4.10b-d). As the viscosity increases from 0.5 to 1.5 Pa.s, there seems to be also a difference when segmentation contractions were produced at 2 and 3s (Figures 4.10c and 4.10d). These results suggested that maybe the rheological behaviour of CMC at higher frequencies changed somehow that do not represent a physical barrier in mass transfer processes.
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Figure 4.10 Glucose absorption in the SIM for solutions added with CMC at various concentrations (a) 0.25, (b) 0.50, (c) 1.00 and (d) 1.25% (w/v) in a glucose solution (1%, w/v) under segmentation contractions at 1, 2 and 3 seconds.

4.3.2 Overall mass transfer coefficients (OMTC): effect of frequency of segmentation

The OMTC were estimated as described in section 4.2.2 by dividing the rate of absorption obtained from the slope of the curves in Figures 4.8, 4.9 and 4.10 for guar, pectin and CMC, respectively, by the surface area (0.5 m$^2$), $K$ the overall mass transfer coefficient, can hence be calculated from Equation 4.2.
4.3.2.1 Overall mass transfer coefficients for guar

In Table 4.9 the results of the OMTC as a function of both frequency and concentration of guar gum are summarized.

Table 4.9 OMTC for guar gum solutions as a function of mixing frequency.

<table>
<thead>
<tr>
<th>Concentration of guar gum (%, w/v)</th>
<th>1s ((10^{-7}\text{ m/s}))</th>
<th>2s ((10^{-7}\text{ m/s}))</th>
<th>3s ((10^{-7}\text{ m/s}))</th>
</tr>
</thead>
<tbody>
<tr>
<td>0.00</td>
<td>4.57 ± 0.10</td>
<td>4.56 ± 0.11</td>
<td>4.36 ± 0.22</td>
</tr>
<tr>
<td>0.25</td>
<td>4.48 ± 0.12</td>
<td>4.65 ± 0.02</td>
<td>4.66 ± 0.39</td>
</tr>
<tr>
<td>0.50</td>
<td>3.53 ± 0.06</td>
<td>3.55 ± 0.23</td>
<td>3.62 ± 0.02</td>
</tr>
<tr>
<td>0.65</td>
<td>3.33 ± 0.06</td>
<td>3.35 ± 0.09</td>
<td>3.47 ± 0.05</td>
</tr>
<tr>
<td>0.75</td>
<td>3.30 ± 0.20</td>
<td>3.37 ± 0.15</td>
<td>3.43 ± 0.01</td>
</tr>
</tbody>
</table>

The OMTC estimated for guar was not affected by the mixing frequencies used in these experiments. Although a significant reduction of the OMTC was observed when the concentration of guar was increased from 0.25 to 0.50% (w/v), for all the frequencies used, increasing the concentration of guar from 0.5 to 0.75% (w/v) did not appear to have a significant impact on the OMTC. Figure 4.11 shows the OMTC for guar as a function of the number of contractions (Figure 4.11a) and viscosity of the solution (Figure 4.11b). Clearly, mass transport was reduced up to 30% as the viscosity increased from 0.02 to 0.42 Pa.s. Nevertheless, it seems that no higher reduction was reached for guar-containing solutions with zero shear viscosities of 1.2 and 3.2 Pa.s.
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Results show that the OMTC falls from $4.6 \times 10^{-7}$ m/s for diluted solutions with low zero-shear viscosities (0.02 Pa.s) to up to $3.3 \times 10^{-7}$ m/s for high-viscosity solutions (3.2 Pa.s). Overall, food formulation specifically viscosity reduced mass transfer processes when guar gum was added to solutions of glucose independently of the frequency of mixing applied to the system. As expected, OMTC were higher for low viscosity solutions (water-like viscosity). Nevertheless, when the viscosity of the solution increases from 0.4 to 3.2 Pa.s, there was no further effect on reduction of the OMTC. Results suggest that shear rates in the system can be in the range 10 to 100 (1/s) at infinite viscosity (Figure 3.5a), where no significant differences were observed for viscosity of guar solutions.

4.3.2.2 Overall mass transfer coefficients for pectin

From Table 4.10, it can be seen that values of the OMTC for pectin were similar at all three frequencies as found for guar. Although the value of the OMTC tends to increased as the “mixing frequency” was increased no significant difference was observed. Like guar, pectin is a polymer that has been used in diets for diabetic people and it has proved to be effective in reducing glucose levels in blood. As expected, the change in viscosity was driven to a
reduction of the OMTC. Clearly, by increasing the concentration of pectin the OMTC dropped drastically for all frequencies applied. When the concentration of pectin increased by twofold (2.0%, w/v), the OMTC was reduced about 20% at 1s and up to 30% when contractions were applied every 2s to the system.

Table 4.10 OMTC for pectin solutions as a function of mixing frequency.

<table>
<thead>
<tr>
<th>Concentration of pectin (%, w/v)</th>
<th>1s</th>
<th>2s</th>
<th>3s</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>(10^{-7} m/s)</td>
<td>(10^{-7} m/s)</td>
<td>(10^{-7} m/s)</td>
</tr>
<tr>
<td>1.0</td>
<td>3.77 ± 0.14</td>
<td>3.89 ± 0.13</td>
<td>---</td>
</tr>
<tr>
<td>2.0</td>
<td>3.31 ± 0.11</td>
<td>3.37 ± 0.24</td>
<td>3.44 ± 0.26</td>
</tr>
<tr>
<td>2.5</td>
<td>3.13 ± 0.02</td>
<td>3.15 ± 0.02</td>
<td>3.30 ± 0.06</td>
</tr>
<tr>
<td>3.0</td>
<td>2.98 ± 0.20</td>
<td>3.00 ± 0.06</td>
<td>3.11 ± 0.03</td>
</tr>
</tbody>
</table>

Figures 4.12 (a) and (b) show the OMTC as a function of both segmentation contractions and food formulation (viscosity). Once more, the mixing frequency did not appear to affect significantly glucose absorption when pectin was added to the solution. A clear difference in mass transport was only observed between dilute solutions (1.0%, w/v) and solutions with concentration of pectin 2.0% (w/v). The OMTC values were clearly reduced by 30% as the viscosity of the solution increased.
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Figure 4.12 OMTC for glucose-pectin solutions as a function of (a) the number of contractions per minute applied to the SIM and (b) zero shear viscosity of the solution.

4.2.3.3 Overall mass transfer coefficients for CMC

The results of the OMTC as a function of both frequency and concentration of CMC are presented in Table 4.11. As shown previously for absorption, the frequency of the segmentation contractions had a great impact on mass transfer when CMC was used. On the other hand, increasing the viscosity of the digesta by adding CMC slightly reduced the OMTC. At 2s for example, the OMTC was reduced only 20% when the viscosity was increased ten-fold. However, when the viscosity increased by three times no significant difference was observed for the OMTC.

Table 4.11 OMTC for CMC solutions as a function of frequency of segmentation

<table>
<thead>
<tr>
<th>Concentration of CMC (% w/v)</th>
<th>1s ( (10^{-7} \text{ m/s}) )</th>
<th>2s ( (10^{-7} \text{ m/s}) )</th>
<th>3s ( (10^{-7} \text{ m/s}) )</th>
</tr>
</thead>
<tbody>
<tr>
<td>0.25</td>
<td>4.19 ± 0.09</td>
<td>4.24 ± 0.01</td>
<td>----</td>
</tr>
<tr>
<td>0.50</td>
<td>1.49 ± 0.34</td>
<td>3.79 ± 0.29</td>
<td>3.87 ± 0.18</td>
</tr>
<tr>
<td>1.00</td>
<td>1.55 ± 0.05</td>
<td>3.06 ± 0.13</td>
<td>3.31 ± 0.01</td>
</tr>
<tr>
<td>1.25</td>
<td>1.08 ± 0.15</td>
<td>3.05 ± 0.04</td>
<td>3.24 ± 0.04</td>
</tr>
</tbody>
</table>
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Unlike guar and pectin, the OMTC estimated from the CMC-containing solutions were significantly influence by the number of segmentation contractions, decreasing up to threefold when the frequency was increased from 7 - 10 to 20 contractions per minute.

Figure 4.13 (a) and (b) show the OMTC as a function of both segmentation contractions and food formulation. The mixing frequency appears to affect significantly glucose absorption when CMC was added to the solution.

Figure 4.13 OMTC for glucose-CMC solutions as a function of (a) the number of contractions per minute applied to the SIM and (b) zero shear viscosity of the solution.

Figure 4.13 (a) and (b) shows that for low viscosities (0.08 Pa.s) solution of CMC the effect of frequency was not significant, with similar OMTC. However, when the viscosity of the lumen increased the mass transfer and the mixing frequency have a significant effect on the mass transfer phenomena. When the number of contractions applied to the SIM increases from 7 to 10 pulses/min, the OMTC decreased (Figure 4.13a). Increasing the frequency of the contractions may well decrease the amplitude of the fluid pulsing along the SIM, and therefore mass transfer decreases when pulses were applied every second. Overall, for glucose solution supplemented with CMC, the frequency of segmentation contractions has a

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significant effect (P<0.001) on the mass transfer coefficient. As segmentation decreases from 1 to 3s the OMTC increased.

Diffusion to the membrane interface is one mechanism for mass transfer, however, small volumes or elements of this interfacial region are not static, but are constantly exchanged or replaced with new elements from the bulk region (Cussler, 2000). ‘Surface renewal’ theories or mass transfer describe the process in terms the length of time that small fluid elements spend in the interfacial area. The influence of the wall movements on the transfer process is significant, therefore.

4.3.3 Local mass transfer coefficients

4.3.3.1 Local mass transfer coefficient for guar gum

The range of mass transfer coefficients in the lumen side (biopolymer side) are shown in Figure 4.14 when segmentation movements were applied.

In accordance with the results for the OMTC, as the viscosity of the digesta was raised from 0.02 to 0.4 Pa.s by increasing the guar concentration, the resistance of the biopolymer to mass transfer substantially increases reaching a plateau for viscosities higher than 1.0 Pa.s. As such there is a clear reduction in the biopolymer side mass transfer coefficient as a consequence of increasing concentrations of guar gum up to 0.75% (w/v).
Figure 4.14 Local mass transfer coefficients of the biopolymer side from different glucose guar solutions.

The oscillatory Reynolds number of the fluid in the biopolymer side was calculated by equation 4.6. The maximum velocities were estimated by using the volume displaced by segmentation contractions produced at 1, 2 and 3 seconds, respectively, data are shown in Table 4.12. These velocities were calculated by assuming that the volume the cuff displaced during each contraction is equal to $9.65 \times 10^{-5} \text{ m}^3$ as previously described in section 4.2.3.

<table>
<thead>
<tr>
<th>Time (s)</th>
<th>Volumetric flow rates ($10^{-5} \text{ m}^3/\text{s}$)</th>
<th>Mean velocity (m/s)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>9.65</td>
<td>0.1365</td>
</tr>
<tr>
<td>2</td>
<td>4.82</td>
<td>0.0680</td>
</tr>
<tr>
<td>3</td>
<td>3.21</td>
<td>0.0455</td>
</tr>
</tbody>
</table>
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Based on these data and the properties of guar solutions, oscillatory Reynolds and Sherwood numbers were estimated for each mixing frequency and results are presented in Table 4.13.

Table 4.13 Oscillatory Reynolds numbers and Sherwood estimated for guar solutions.

<table>
<thead>
<tr>
<th>Zero shear viscosity (Pa.s)</th>
<th>1s Re₀</th>
<th>1s Sh₀</th>
<th>2s Re₀</th>
<th>2s Sh₀</th>
<th>3s Re₀</th>
<th>3s Sh₀</th>
</tr>
</thead>
<tbody>
<tr>
<td>0.02</td>
<td>1040</td>
<td>996</td>
<td>2076</td>
<td>1161</td>
<td>3113</td>
<td>1047</td>
</tr>
<tr>
<td>0.42</td>
<td>56</td>
<td>69</td>
<td>112</td>
<td>70</td>
<td>168</td>
<td>77</td>
</tr>
<tr>
<td>1.20</td>
<td>19</td>
<td>55</td>
<td>38</td>
<td>55</td>
<td>57</td>
<td>60</td>
</tr>
<tr>
<td>3.19</td>
<td>6</td>
<td>55</td>
<td>12</td>
<td>57</td>
<td>18</td>
<td>61</td>
</tr>
</tbody>
</table>

Results show that as the viscosity of the glucose-guar solution increased, the Re₀ also decreased. Low-viscosity solutions (0.02 Pa.s) driven to oscillatory Reynolds numbers in the transition and turbulent flow region in which turbulent mixing occurs enhancing mass transfer processes. In addition, Sherwood number was inversely correlated to the guar concentration decreasing as the viscosity increased, pointing out that mass transfer processes occur predominately by diffusion for viscous solutions. High Reynolds and Sherwood numbers (Figure 4.15) were obtained as the frequency of contractions decreased from 20 to 7 contractions per minute produced every 1 or 3 seconds, respectively. These results suggested that the higher the intensity of the contractions is the deeper the contraction i.e. major volume is displaced improving the mixing of the lumen contents.
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Figure 4.15 Sherwood versus oscillatory Reynolds numbers for guar solutions at different mixing frequencies (1, 2, and 3 seconds).

4.3.3.2 Local mass transfer coefficient for pectin solutions

The mass transfer coefficients in the lumen side (biopolymer side) for pectin are shown in Figure 4.16 when segmentation movements were applied.

As the viscosity of the digesta was raised from 0.05 to 2.00 Pa.s (Figure 4.12), by increasing the concentration of pectin, the biopolymer mass transfer coefficient \( k_{bp} \) substantially decreases. As such there is a clear reduction in the biopolymer side mass transfer coefficient as a consequence of increasing concentrations of pectin up to 3.0% (w/v).
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Figure 4.16 Local mass transfer coefficients of the biopolymer side from different glucose pectin solutions.

Oscillatory Reynolds and Sherwood number were estimated for each mixing frequency and results are presented in Table 4.14. Again, Sherwood number was inversely correlated to the pectin concentration, decreasing as the viscosity increased. Reynolds in the transition region and Sherwood numbers were relatively higher as the frequency increased suggesting that at 3s segmentation contractions displace major volume improving the mixing of the lumen contents in the SIM (Figure 4.17).

Table 4.14 Oscillatory Reynolds numbers and Sherwood estimated for pectin solutions.

<table>
<thead>
<tr>
<th>Zero shear viscosity (Pa.s)</th>
<th>1s</th>
<th>2s</th>
<th>3s</th>
</tr>
</thead>
<tbody>
<tr>
<td>Re₀</td>
<td>Sh₀</td>
<td>Re₀</td>
<td>Sh₀</td>
</tr>
<tr>
<td>0.05</td>
<td>532</td>
<td>96</td>
<td>1063</td>
</tr>
<tr>
<td>0.25</td>
<td>106</td>
<td>53</td>
<td>213</td>
</tr>
<tr>
<td>0.70</td>
<td>37</td>
<td>44</td>
<td>75</td>
</tr>
<tr>
<td>2.00</td>
<td>14</td>
<td>38</td>
<td>28</td>
</tr>
</tbody>
</table>
Figure 4.17 Sherwood versus oscillatory Reynolds numbers for pectin solutions at different mixing frequencies (1, 2, and 3 seconds).

4.3.3.3 Local mass transfer coefficient for CMC solutions

The local mass transfer coefficients (biopolymer side) for CMC solutions are shown in Figure 4.18 when segmentation movements were applied.

As the viscosity of the digesta was raised from 0.08 to 1.45 Pa.s, by increasing the concentration of CMC, the biopolymer mass transfer coefficient ($k_{bp}$) substantially decreases. There is a clear reduction in the biopolymer side mass transfer coefficient as a consequence of increasing concentrations of CMC up to 1.25% (w/v).
Based on data from $k_{bp}$ for CMC, oscillatory Reynolds and Sherwood number were estimated for each frequency and results are presented in Table 4.15. For CMC Sherwood number was inversely correlated to the concentration of biopolymer used, decreasing as the viscosity increased. Reynolds were in the transition region and Sherwood numbers were relatively higher as the frequency increased suggesting that at 3s segmentation contractions displace major volume improving the mixing of the lumen contents in the SIM (Figure 4.19).
Table 4.15 Oscillatory Reynolds numbers and Sherwood estimated for CMC solutions.

<table>
<thead>
<tr>
<th>Zero shear viscosity (Pa.s)</th>
<th>1s</th>
<th>2s</th>
<th>3s</th>
</tr>
</thead>
<tbody>
<tr>
<td>0.09</td>
<td>270</td>
<td>229</td>
<td>540</td>
</tr>
<tr>
<td>0.30</td>
<td>115</td>
<td>10</td>
<td>231</td>
</tr>
<tr>
<td>0.50</td>
<td>43</td>
<td>10</td>
<td>87</td>
</tr>
<tr>
<td>1.45</td>
<td>16</td>
<td>17</td>
<td>32</td>
</tr>
</tbody>
</table>

Figure 4.19 Sherwood versus oscillatory Reynolds numbers for CMC solutions at different mixing frequencies (1, 2, and 3 seconds).

The comparison of absorption of glucose with open or close SIM configurations disclosed highly significant differences. The absorption rate was higher when recirculation of the chyme was done (open-SIM) with up to 50% of increasing of the absorption rate.

The results indicate that, as a whole, the absorption rates i.e. OMTC were higher when the viscosity of the food increased.
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As the number of contractions per minute increases, glucose absorption decreases (Figure 4.19). For low viscosity, changes in the number of pulses per minute does not have a significant impact on glucose absorption having only about 10% of absorption after one hour. However, as the viscosity increases absorption of glucose was clearly modified by frequency.

As shown by Macagno et al. (1982), when different types of contractions are applied different degrees of convective transport are induced because of the different kinematics and volumes that they displaced.

- ‘1s’ generates 20 pulses per minute this is a very short contraction that displace a very small volume and less longitudinal mixing.
- ‘2s’ is equal to 10 pulses per minute this is a deeper contraction compare with the one generated at 1s, with a major volume displace.
- At 3s, the deepest contraction was generated displacing the highest volumes through the SIM and as a consequence convective transport in the lumen side was achieved.

As the frequency increases but the time of exposure to diffusion decreases the absorption of glucose decreases as well. However for 3s the time of exposure increases driven to higher glucose absorption enhancing mass transfer.

The effect of the type of polymer and its rheological behavior at characteristic physiological shear rates must be taken into account to predict the effect that biopolymers such as guar, pectin or CMC has on glucose or actives absorption.

Each contraction reduces the area available for absorption occluding the flexible tube- lumen and recipient sides- leaving no much area for absorption. Absorption has place when the tubing is relaxed.
These experiments suggest that glucose absorption in solution containing guar and pectin in the range of concentration tested is little affected by the mixing frequencies used in this work.

In contrast, CMC showed to have a significant difference among processing conditions. This could be explained based not only on the rheological behaviour of the biopolymer employed but also based on the chemical structure and composition of this biopolymer compared to guar (discussed in next section). According to the viscosity and the results of the OMTC in which no great difference was observed at zero shear viscosities higher than 0.1 Pa.s. It can be suggested the working shear rate could be from 10 to 50-100 (1/s) at which guar and pectin seems to have similar viscosity independently of the concentration of biopolymer used. On the other hand, CMC has different viscosities at this range of shear rates this could explain why only with CMC significant differences were observed at carious frequencies.

Reduction on absorption of glucose in the presence of guar is probably due to resistance by viscous solutions of convective effect of the intestinal contractions.

In the GI tract the frequency, amplitude and duration of segmentation contractions depend on the physical and chemical properties of the chyme. Large meals, for example, stimulate powerful phasic contractions through the body-antral reflex in the stomach (Schulze, 2006). Liquid foods stimulate deep contractions whereas shallow contractions are generated when contents are highly viscous (Schulze, 2006). Propagative and non propagative contractions generate flow effects that are likely to enhance digestion and absorption. It is believed that intestinal motility in vivo increases the exposure of the mucosa to absorbable materials.

The concentration gradients of any active in chyme that is being absorbed could be modified by segmentation as a result of the dispersion, spread, mixing transport and distribution of particles present in the chyme. By understanding these phenomena it will be possible to
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influence on the design of structured food and drug delivery systems to provide specific benefits to consumers.

4.4. Biopolymer and their impact on absorption

Within the range of soluble fibres used in this work, the OMTC were reduced as the concentration/viscosity of the biopolymer increase. Each polymer acts as a physical barrier to nutrient absorption contributing to the formation of unstirred layer adjacent to the mucosa (Anttila et al., 2003). However, each biopolymer shows different effect on glucose absorption that could be related to their rheological behaviour.

4.4.1. Absorption versus rheology of the biopolymers

While pectin has a Newtonian behaviour in most of the shear rates used (0.1 – 1000 s⁻¹), guar gum and CMC are non-Newtonian/pseudoplastic fluids with shear thinning characteristics i.e. viscosity decreases as shear rate increases (see Figure 3.5). To understand how their rheological behaviour affects absorption, the OMTC of each biopolymer (section 4.3.2) are compared in the following section. Only three cases of study were selected:

(i) Biopolymers (guar, pectin and CMC) at the highest concentration

(iii) Biopolymers (pectin and CMC) with similar zero-shear viscosities

(iv) Biopolymers (guar and pectin) with a cross over at shear rate of 100 (1/s)

Case I. A comparison among viscometry profiles of guar (0.75%, w/v), pectin (3.0%, w/v) and CMC (1.25%, w/v) is presented in Figure 4.20. Table 4.16 shows the values of the OMTC for each biopolymer at the highest concentration used.
Table 4.16 OMTC for guar, CMC and pectin are compared at the highest concentration used.

<table>
<thead>
<tr>
<th>Biopolymer</th>
<th>2s</th>
<th>3s</th>
</tr>
</thead>
<tbody>
<tr>
<td>Guar (0.75%, w/v)</td>
<td>3.37 ± 0.15</td>
<td>3.43 ± 0.01</td>
</tr>
<tr>
<td>CMC (1.25%, w/v)</td>
<td>3.05 ± 0.04</td>
<td>3.24 ± 0.04</td>
</tr>
<tr>
<td>Pectin (3.0%, w/v)</td>
<td>3.00 ± 0.06</td>
<td>3.11 ± 0.03</td>
</tr>
</tbody>
</table>

**Figure 4.20** Rheological behaviour of guar (0.75%, w/v), pectin (3.0%, w/v) and CMC (1.25, w/v).

As shown in Figure 4.20 guar gum shows the highest low-shear viscosities however is also strongly shear-thinning. However, CMC and pectin have similar behaviours at low shear rates (0.1 to 50 s⁻¹). The three polymers have a crossover at shear rate of ~50 s⁻¹. At high shear rates 100 - 1000 s⁻¹, this trend changed and pectin has the highest viscosity, then CMC and finally guar. Interestingly, the OMTC at low frequencies of segmentation 2 and 3s (Table 4.16) are in agreement with the rheological behaviour of the biopolymers at high shear rates. In this case, the OMTC for pectin were lower than CMC and guar. Based on the OMTC, pectin was more effective than CMC to reduce absorption, and CMC was more effective than guar.
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Overall, guar gum with the highest zero shear viscosity value displayed the highest OMTC (Table 4.16), i.e. glucose absorption. However, pectin and CMC with similar zero shear viscosities had similar OMCT values at 2s frequency. Results from Table 4.16 suggested that pectin was the most effective biopolymer on reducing glucose absorption. This could be explained by the rheological behaviour of the biopolymers at high shear rates after 100 s\(^{-1}\) where pectin has the highest viscosity, then CMC and finally guar.

**Case II.** It was also possible to observe that similar zero shear viscosities from pectin and CMC resulting in similar OMTC (Table 4.17 and Figure 4.21).

Table 4.17 OMTC for CMC and pectin are compared at different concentration.

<table>
<thead>
<tr>
<th>Biopolymer</th>
<th>2s</th>
<th>3s</th>
</tr>
</thead>
<tbody>
<tr>
<td>Pectin (2.5%, w/v)</td>
<td>3.15 ± 0.02</td>
<td>3.30 ± 0.06</td>
</tr>
<tr>
<td>CMC (1.0%, w/v)</td>
<td>3.06 ± 0.13</td>
<td>3.31 ± 0.01</td>
</tr>
</tbody>
</table>

![Figure 4.21](image.png)  
**Figure 4.21** Rheological behaviour of pectin (2.5%, w/v) and CMC (1.0%, w/v).
Case III. On the other hand, when the viscometry profiles from guar and pectin (Figure 4.22) showed a crossover at 100 s\(^{-1}\) the OMTC (Table 4.18) were very similar. These figures confirm that the working shear rates in the SIM could be considered around 100 s\(^{-1}\).

Table 4.18 OMTC for guar and pectin are compared at different concentrations.

<table>
<thead>
<tr>
<th>Biopolymer</th>
<th>1s</th>
<th>2s</th>
<th>3s</th>
</tr>
</thead>
<tbody>
<tr>
<td>Guar (0.65%, w/v)</td>
<td>3.33 ± 0.06</td>
<td>3.35 ± 0.09</td>
<td>3.47 ± 0.05</td>
</tr>
<tr>
<td>Pectin (2.0%, w/v)</td>
<td>3.31 ± 0.11</td>
<td>3.37 ± 0.24</td>
<td>3.44 ± 0.26</td>
</tr>
</tbody>
</table>

Figure 4.22 Rheological behaviour of guar (0.625%, w/v) and pectin (2.0%, w/v).

Overall, the different flow characteristics of each biopolymer are of practical value in determining the type of gum required to delay glucose uptake form the intestinal lumen. However, in some cases rheological behaviour is limited and the results could be controversial and not conclusive. For that reason, it is fundamental to understand how chemical structure of each biopolymer affects the interactions among chains (entanglements) that could also influence biopolymers networks structure in lumen and most importantly in
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the stagnant region or the unstirred layer in the gut. Both physical and chemical interactions are important to the physiological functions of the polymer in the GI tract.

4.4.2 Absorption and chemical structure of the chyme

4.4.2.1 Guar gum and chemical structure

Guar gum is a neutral biopolymer consisting of 1,4-linked β-D-mannopyranose back bone with the branches of 1,6-linked α-D-galactopyranose (Figure 4.23). Previous studies in vivo have stated that comparing the three polymers used in this work on postprandial glucose and insulin levels, guar showed to be the most effective biopolymer on reducing plasma glucose levels (Brenelli et al., 1997). This behaviour was explained as guar showed the greatest stability and viscosity under acidic conditions in vitro i.e. under gastric conditions. Being non-ionic, guar gum is not significantly affected by ionic strength or pH. Although guar results in slightly higher OMTC, the concentration of guar used to increased the viscosity of the digesta was ten-fold less than pectin. This criterion must be also considered in selecting ingredients to develop functional food or drug formulations.

Figure 4.23 A schematic of the chemical structure of guar gum (Chaplin, 2012)
4.4.2.2. Pectin and chemical structure

On the other hand, pectins are rich in galacturonic acid groups with linear chains (Figure 4.24). Pectins are divided into two major groups on the basis of their degree of esterification that result in different functional properties.

![Figure 4.24 A schematic of chemical structure of pectin (Chaplin, 2012)](image)

The association of pectin chains leads to the formation of tridimensional networks that leads to gel formation in presence of divalent ions such as Ca$^{++}$. In human digestion, studies have revealed that pectin could bind to cholesterol in the GI tract slowing absorption by entrapping carbohydrates (Kim, 2005). Therefore, high-methoxy pectin showed to be more effective than low-methoxy pectin on reducing blood glucose levels (Kim, 2005).

Polysaccharides with some charged groups such as pectins will be generally soluble in saline solution due to the electrostatic repulsion that inhibits ordered packing (Guillon and Champ, 2000).

4.4.2.3 CMC and chemical structure

On the other hand, CMC has carboxymethyl groups bound to the hydroxyl groups of the glucopyranose structure (Figure 4.25). As guar, solutions of sodium-CMC are non-Newtonian with a pseudoplastic behaviour. Degree of pseudoplasticity results from the nonuniform substitution of the carboxymethyl groups along the cellulose chain. The
properties of sodium-CMC can be controlled by varying the uniformity of substitution and the degree of polymerization.

![Figure 4.25](image.png) A schematic of the chemical structure of the CMC (Chaplin, 2012)

In this study highly viscous CMC shows to be more effective on reducing glucose absorption at high frequencies of segmentation (1s). This could be correlated to its rheological behaviour at high shear rates (50-100 s$^{-1}$) and its chemical structure that possible formed strong interactions among its chains i.e. a denser network was formed.

Overall, the influence that viscosity of the chyme has on postprandial glycemia has been assessed in a variety of studies; results *in vivo* are continued being controversial. Thus, rheological studies and chemical evaluation must be combined to predict their effects on cholesterol and glucose blood levels. For example, information on molecular weight, concentration and purity of the soluble fibres most be considered for selection of food ingredients to obtain best end-results.

### 4.5 Conclusions

In summary, mass transfer in the SIM was enhanced by up to 30% when segmentation was applied to the system, reducing the unstirred layer formed near the membrane helping the
diffusion of glucose. Mixing processes increase the rate of glucose transferred by bringing material to the epithelium. In contrast, mixing was impaired by the viscous fluids used: guar and CMC.

When the viscosity of the lumen increases the effect of mixing was reduced by up to 90% for 0.5% (w/v) CMC solutions (no significant difference was observed between segmentation and no segmentation for viscous solutions).

Overall, mixing and food formulation of the sample have a significant effect on the mass transfer phenomena of glucose absorption in the SIM. Whilst mixing enhances mass transfer phenomena in the intestine, increasing the viscosity of the food results in a reduction of the mass transfer across the membrane of the SIM.

Wall movements influence intestinal absorption by bringing material to the sites of absorption. However, the only biopolymer that seems to have a significant effect on glucose absorption at different frequencies was CMC. Mass transfer coefficients were very similar when guar gum and pectin were added to the glucose solution, as a function of different frequencies of contractions.

Overall, results show that mass transfer coefficients increased as the frequency of contractions decreased. On the other hand, increasing the viscosity of the lumen or biopolymer side resulted in a decrease of the mass transfer coefficients.
CHAPTER 5 - STARCH DIGESTION IN THE SIM

5.1 Introduction

Most of food digestion takes place in the small intestine in which intestinal motility promotes interactions between food particles and digestive fluids. In order to understand and reproduce these processes, an *in vitro* dynamic small intestine model (SIM) was used to study starch digestion. Starch was selected as model in this work due to its importance as glycemic carbohydrate component in foods; contributing significantly to the exogenous supply of glucose and the total food energy intake (Slaughter et al., 2001). This work focuses on the effect that mixing and food formulation has on the rates of starch digestion.

Starch in the human diet is digested in the mouth, stomach and small intestine into di- and monosaccharides, which are the only form which can be absorbed (Guyton and Hall, 2006). According to the rate of hydrolysis in the gut, starch could be classified into rapidly digestible starch (RDS), slowly digestible starch (SDS), and resistant starch (RS). RDS is absorbed in the duodenum and proximal regions of the small intestine within the first 20 minutes, and it is correlated to the glycemic values of the sample. On the other hand, RS is the starch not hydrolysed after 120 minutes of incubation, i.e. not digested in the upper gastrointestinal tract, while SDS is the digested portion between RDS and RS and it is thought to be digested along the whole small intestine between 20 and 120 minutes (Englyst et al., 1992, van Kempen et al., 2010). Concerning to glucose delivery, RDS provides a rapid and high level of glucose (high glycaemic index), RS theoretically does not provide glucose and SDS provides a slow and prolonged glucose delivery (low glycaemic index). Control of glucose delivery in foods can be determined by food structure and the physicochemical properties of the carbohydrate components in foods (Lundin et al., 2008, Snow and O'Dea, 1981). When
refined sugars and rapidly digested carbohydrates are included on daily diets, blood glucose and insulin responses are usually enhanced, resulting in dietary related chronic diseases such as diabetes and obesity (Read and Welch, 1985). Since the relationship between an ingested carbohydrate and its related glycaemic index value is not always intuitively obvious (Monell, 2005), structured foods with specific health benefits should be designed in conjunction with studies to understand how food formulation impacts on digestion and absorption rates in the gastrointestinal tract.

As established in section 4.2, intestinal absorption was influenced by the physical properties (viscosity, density, concentration, flow rate) of the lumen content. Increasing viscosity of the lumen reduces the rate of nutrient absorption and mass transfer to the membrane. On this context, addition of viscous fibres into foods may also protect starch from enzymatic attack (Leclere et al., 1994). Such as rates of digestion and nutrients absorption could be modified by changing food formulation resulting in structured food with improved health properties.

The main objective of the experiments presented in this Chapter was to reproduce and model starch digestion by studying the effect that mixing and food formulation has on glucose absorption. Furthermore, starch digestion was studied varying the concentration of starch, α-amylase and flow rate at which the enzyme solution was fed. Digestion and absorption processes were studied in the SIM by simulating as close as possible the physiological conditions of the human small intestine. Two different SIM configurations were used: (i) closed-SIM, with no recirculation of the chyme and (ii) open-SIM, in which recirculation of the chyme was induced by a peristaltic pump.
5.2 Starch hydrolysis in the closed SIM configuration

5.2.1 Starch digestion: effect of mixing

Due to the importance of mixing on enzymatic digestion processes, in this section, starch hydrolysis in the SIM is compared with the hydrolysis performed in a well-mixed reactor as described in section 3.4.5.

Model I. The Stirred Tank Reactor (STR) was equipped with a Rushton turbine impeller and four baffles as described in section 3.4.5.4 (Figure 3.7a). The diameter of the tank and the impeller were 0.075 m, and 0.045 m, respectively. The turbine was operated at 300 rpm and reactions were carried out in 250 ml of aqueous solutions containing 10 g/l of corn starch pregelatinized during 10 minutes in boiling water.

Model II. In the Small Intestine Model (SIM) mixing was induced by two inflatable cuffs (Figure 3.7b) to represent the characteristic segmentation contractions in the small intestine. Contraction were generated by two inflatable cuffs at a rate of ten cycles per minute with the following standard pattern: 2 seconds inflation, 2 seconds deflation and 2 seconds delay between each cuff. Reactions were carried out in the lumen side (membrane side) of the SIM with 600 ml of aqueous solutions containing 10 g/l of corn starch previously gelatinized (10 minutes in boiling water) and with no recirculation of the chyme.

Enzyme solution (25 ml for the STR and 50 ml for the SIM) was freshly prepared for the digestion analysis by dissolving the pancreatic α-amylase (5 U/ml) and amyloglucosidase (0.5 U/ml) into a pancreatic mix solution for 10 minutes at room temperature (according to the method described in section 3.4.5.2). According to the supplier, one unit of pancreatic amylase will liberate 1.0 mg of maltose from starch in 3 min at pH 6.9 at 20 °C.
Chapter 5 - Starch digestion

The pancreatic solution was fed into the reactor using a thin polyvinylchloride tube ending approximately 5.0 cm distally to the first cuff of the SIM (Figure 5.1) and 3.0 cm from the base of the STR. Before hydrolysis, the tubing was connected to a micro peristaltic pump and filled with the pancreatic solution. This micro-system was implemented here to mimic the injection of enzymes and salts from the pancreas into the small intestine. The pancreatic duct (see section 2.2.6) was represented by the long tubing (0.003 m of diameter) showed in Figure 5.1.

![Figure 5.1 Schematic of the experimental SIM set-up for starch hydrolysis.](image)

The reaction was started by pumping the pancreatic mix solution into the reactor of the STR or in the lumen side of the SIM at an apparent flow rate of 3.0 ml/min (Table 5.1). At time zero before injection of the pancreatic solution, a sample was taken and used as control/blank for glucose determination.

Table 5.1 Experimental conditions used for starch digestion.

<table>
<thead>
<tr>
<th></th>
<th>Starch</th>
<th>Enzyme</th>
<th>Flow rate</th>
</tr>
</thead>
<tbody>
<tr>
<td>STR</td>
<td>1.0%, w/v</td>
<td>5 U/ml</td>
<td>3.0 ml/min</td>
</tr>
<tr>
<td>SIM</td>
<td>1.0%, w/v</td>
<td>5 U/ml</td>
<td>3.0 ml/min</td>
</tr>
</tbody>
</table>
Samples were taken from the STR and the SIM reaction mixture at time intervals up to 90 minutes before determination of reducing sugars. Since the closed-SIM configuration was used, samples from the lumen side of the SIM were taken with a needle and syringe placed in the peristaltic pump tubing, at the distal end of the membrane. To inactivate the enzyme the glass tube containing the sample was placed into boiling water during 5 minutes. The concentration of glucose was calculated following the method of 3,5-dinitrosalicylic acid (DNS) for reducing sugars (expressed as ‘mM’ of glucose). All experiments were performed at room temperature and in triplicate.

Figure 5.2 shows the increase in glucose concentration after starch hydrolysis in both models the STR and the SIM. By using the STR, starch digestion has a fast rate of reaction (44.8 µM/s) at the beginning of the hydrolysis process following by a decreased rate, reaching a plateau (saturation of the enzyme) after approximately 20 min with maximum glucose concentration of 31 mM. On the other hand, the hydrolysis of starch in the SIM appears to have a sigmoidal behavior with a ‘lag phase” due to the incomplete mixing of the starch and amylase at the early stage of the hydrolysis within the first ten minutes. After the lag time, kinetics of starch in the SIM appeared to have the same trend than those in the STR, with a rapid rate (24 µM/s) from 10 to 30 minutes followed by a decreasing rate until equilibrium is reached. Maximum glucose concentration (31 mM) in the SIM was achieved after 40 minutes of digestion. This behaviour depends on the mixing mechanism applied to the system. Faster reactions took place in the STR, in which enzymes and substrate were well-mixed, due to the turbulence created by the baffles and the impeller at 300 rpm with Re = 11300. On the contrary, fluids in the SIM are in the transitional regime (Re = 2100) in which mixing was provided as concentric contractions (10 per minute) by means of the two inflatable rubber
cuffs section. The SIM represents a more realistic model with representative anatomical and physiological features and flow occurring in the human small intestine.

![Time (min)](chart)

![Glucose concentration (mM)](chart)

**Figure 5.2** Starch digestion in the Stirred Tank Reactor (STR) and in the Small Intestine Model (SIM)

The hydrolysis reaction behaviour in the STR and in the SIM was assumed to follow first order reaction in the first minutes of the reaction. Therefore, it is assumed that the reaction follows first order equation which is arranged as follows:

\[
\frac{d[S]}{dt} = -k[S]
\]

(5.1)

Integration of equation 5.1 gives:

\[
\ln \frac{[S]}{[S]_0} = -k t
\]

(5.2)

where \( S \) is the concentration of starch at time \( t \), \( S_0 \) is the concentration of starch (mM) at \( t = 0 \), \( k \) is the kinetic constant (min\(^{-1}\)), and \( t \) is time (Levenspiel, 1999). Based on the data from Figure 5.2, starch concentration was estimated according to equation 5.2. The data were analysed by linear regression where the slope of the curve represents the value of \(-k\) (Table
5.2). Kinetic behaviour trends give a fairly good correlation to first order reaction behaviour marked by the $r^2$ value of 0.94. However, kinetic behaviour of starch hydrolysis in the SIM (Figure 5.2) gives a correlation with $r^2$ less than 0.9. This is a result of the characteristic “lag phase” at the onset of the reaction (first 10 minutes).

Table 5.2 Kinetic constant based on first order reaction behaviour

<table>
<thead>
<tr>
<th>Model</th>
<th>Mixing method</th>
<th>Kinetic constant, $k$ (min$^{-1}$)</th>
</tr>
</thead>
<tbody>
<tr>
<td>STR</td>
<td>Impeller (300 rpm)</td>
<td>0.052</td>
</tr>
<tr>
<td>SIM</td>
<td>Segmentation contractions (2s)</td>
<td>0.012</td>
</tr>
</tbody>
</table>

The results of kinetic parameters described previously provide strong evidence that mixing mechanism has a considerable effect on constant values, thus, it affects the rate of hydrolysis reaction as well. Turbulent flow leads to rapid hydrolysis reaction.

From this experiment, it is possible observed that the type and degree of agitation can affect significantly the rate of starch hydrolysis mostly in the early stage of digestion. During food digestion, mixing is essential to bring substrates and enzymes together and increase the rate of reactions taking place in the GI tract. The small intestine can be modelled as a plug reactor in which the concentration of substrates changes with both time and position.

5.2.2 Starch digestion: effect of food formulation

Corn starch hydrolysis was also studied in the SIM, while varying the viscosity of the digesta by adding guar gum (0.5%, w/v) to a 1.0% (w/v) starch solution and 50 g of white bread (section 3.4.5). For these experiments, glucose concentration was only monitored in the recipient side (outer tube) of the SIM because glucose quantification inside of the membrane is difficult due to viscosity of the digesta Samples (1.0 mL) were taken at time intervals to
estimate glucose absorption. As in section 5.2.1, the concentration of reducing sugars was estimated by the DNS method. All experiments were performed in triplicates.

Figures 5.3 and 5.4 display the change in glucose concentration as a function of time in the recipient side of the SIM, with and without (control) guar added to bread or starch food models. Results showed that glucose absorption was delayed and reduced as the viscosity of the digesta increased. Glucose concentration was reduced by 28% for starch with addition of 0.5% (w/v) guar gum. This result is in agreement with those found previously by Tharakan et al., (2010). Once the reaction was initiated the rate of the starch hydrolysis did not appear to change.

![Graph](image)

**Figure 5.3** Glucose absorption levels in the recipient side of the SIM for 1.0% (w/v) corn starch with and without (control) 0.5% (w/v) of guar gum at 2 s segmentation.

Glucose concentration decreases by 35% for white bread with the same concentration of guar (0.5%, w/v). The evolution of the curve (Figure 5.4) can result in information of the mass transport process taking place. Interestingly for bread, an increasing change in the rate of
absorption illustrated by the crossover in Figure 5.4, was observed. At the beginning of the absorption process, diffusion could be predominant under convection with a lower rate of absorption. However, after approximately one hour, the rate at which glucose and sugars are absorbed was twofold faster; from $0.47 \, \mu\text{M}/\text{s}$ to $0.89 \, \mu\text{M}/\text{s}$. This result could be related to the point at which viscosity of the lumen contents decreases, allowing that glucose mass transport occur faster than in the first stage of absorption likely by convection after glucose generation by starch hydrolysis. In the future, rheology measurements must be done to follow the progress curve of starch hydrolysis and confirm this explanation. As the viscosity of the digesta increases, enzymatic reactions and molecular delivery across the semipermeable membrane operates under mass transfer restriction (Table 5.3.). For rapidly digested foods such as white bread and cooked and homogenised potato, Englyst et al., (1992) pointed out that more than 80% percent of the starch present is converted into glucose within the first 10 minutes. Subsequently, the rate of hydrolysis declines so that only a further 5-10% of the starch is converted to glucose after this period (Englyst et al., 1992).

**Figure 5.4** Glucose absorption levels in the recipient side of the SIM for 50 g of white bread with and without (control) 0.5% (w/v) of guar gum.
This suggests that mass transport to the dialysis membrane could be controlled by the properties of the chyme under physiologically representative process conditions (Tharakan, et al., 2009) and also impact directly on some metabolic disorders such as diabetes and hyperlipidemia. Results are in accordance with previous findings when viscous polymers such as guar, tragacanth and pectin were adding to food systems, decreasing the postprandial glucose and insulin responses (Jenkins et al., 1978).

### 5.2.2.1 Mass transfer coefficients

From data in Figures 5.3 and 5.4, the overall mass transfer coefficient was estimated as described in section 4.2 by the slope in each curve of glucose absorption. The data were analysed in the linear region of the curve, once the flow was fully developed from 10 to 90 min. Absorption appeared to be linear ($R^2$ of 0.95) resulting in a gradient of $7.45 \times 10^{-3}$ mM/s. The flux, $N$, was calculated to be $1.55 \times 10^{-5}$ mol/m$^2$s. By dividing the molar flux by the concentration difference (mmol/m$^3$) an overall mass transfer coefficient, $K$, was obtained as $4.83 \times 10^{-7}$ m/s for the aqueous starch solution and $3.10 \times 10^{-7}$ m/s for the starch solution added with guar, both under segmentation conditions.

The effect of the presence of guar was shown a 36% reduction in the mass transfer coefficient when mixing was applied (Table 5.3). The data not only suggest that the rate of enzymatic reactions was reduced when guar was added but also that viscous solutions can delay the access of actives to the absorptive epithelium due to a decrease of both propulsion and mixing (Read and Welch, 1985). Similarly, this tendency was observed when bread digestion was mimicked in an aqueous solution (control) and in the presence of guar. When viscosity of the lumen content was increased from 2 to 200 mPa.s, glucose absorption was reduced almost 15% after one hour, however, no significant difference was observed within the first minutes of digestion (Figure 5.3 and 5.4) while the overall mass transfer coefficient from bread with
guar gum was reduced 20%. These results are in agreement to those reported by Tharakan et al. (2010).

Table 5.3 Rates of absorption and overall mass transfer coefficient (OMCT) for two food models.

<table>
<thead>
<tr>
<th>Experiment</th>
<th>Absorption (µM/s)</th>
<th>OMTC (µm/s)</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Starch</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Control</td>
<td>0.75 ± 0.05</td>
<td>0.48 ± 0.05</td>
</tr>
<tr>
<td>Guar gum (0.5%, w/v)</td>
<td>0.48 ± 0.02</td>
<td>0.31 ± 0.08</td>
</tr>
<tr>
<td><strong>Bread</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Control</td>
<td>0.67 ± 0.01</td>
<td>0.43 ± 0.04</td>
</tr>
<tr>
<td>Guar gum (0.5%, w/v)</td>
<td>0.50 ± 0.02</td>
<td>0.32 ± 0.02</td>
</tr>
</tbody>
</table>

Overall, mass transfer coefficient was reduced when guar gum (0.5%, w/v) was added to the solution of starch and bread. That could be explained by inhibition of propulsion and mixing in the membrane making interactions between the starch and amylase also difficult. Thus, starch digestion was lower in the presence of guar. Results suggest that molecular delivery is largely influenced by the fluid dynamics of the lumen side. Viscous polymers create a physical resistance for active release and delivery that could modify the rate of nutrients absorption for the design of healthy diets with specific targets. Since diffusion of amylase into the substrate is considered an essential step in carbohydrates hydrolysis mechanisms (Lehmann and Robin, 2007), interactions of starch with fibre, protein and other food components that can prevent effective diffusion and absorption of the enzyme into the starch structure will reduce or delay rates of digestion and further absorption (Collonna et al., 1992; Lehmann and Robin, 2007).

Starch granules were examined by Scanning Electron Microscopy (SEM) before and after hydrolysis. Analysis of the microstructure of the starch was done after one hour of digestion.
in samples of starch previously washed with distilled water and dried in an oven at 50 °C. Scanning electron micrographs of normal corn starch before and after hydrolysis by porcine pancreatic α-amylase are shown in Figures 5.4 (a-d). Before gelatinization and hydrolysis, granules present a spherical-polyhedral shape with a diameter between 2 and 30 µm (Figures 5.4a-b). After cooking and digestion by α-amylase in the presence of guar, formation of clusters was observed due most likely to both cooking and presence guar (Figure 5.4c). Under higher magnification (Figure 5.4d) the starch residue appeared as clustered compactly particles with some fragmented granules (hollow granules) from which the layered granule’s structure could be seen. This clustered structure could also “protect” at some level the granule from amylase delaying enzyme diffusion into the granule of starch.

**Figure 5.5** SEM microphotographs of corn starch (a-b) before hydrolysis and (c-d) after 60 minutes of hydrolysis of pancreatic α-amylase at room temperature in the presence of 0.5% (w/v) of guar gum.
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There is evidence in the literature that enzymatic susceptibility of starch is highly dependable of its microstructure and physicochemical characteristics (Dona et al., 2010). As such any process that can change this orderly structure makes granules of starch available for enzymatic reactions. Digestibility is for example, significantly impacted by shearing and heating the solution of starch (Dona et al., 2010).

5.3. Starch hydrolysis in the open SIM configuration

Having established that mixing and soluble viscous polymers affect digestion and absorption processes in the SIM. A series of experiments were performed to assess the effect that factors such as enzyme activity, flow rate and substrate concentration has of digestion and absorption. However, these experiments were performed in the open-SIM configuration because it was observed that in the closed configuration mixing was not effectively done. As such chyme was recirculated by a peristaltic pump to facilitate mixing and propulsion of the lumen contents. Samples from the lumen side of the SIM were taken from the starch reservoir, no agitation was provided to this vessel to avoid reactions take place there. The volume of the solution in this vessel was kept constant (no more than 150 ml) allowing the volume of liquid displaced by the cuff moves forward and backward. The first set of experiments described in this section was performed to study the effect that enzyme concentration has on glucose absorption. Four experimental conditions were used (Table 5.4), varying (i) the initial concentration of amylase in the pancreatic solution and (ii) the flow rate at which the solution was fed into the lumen side of the SIM. As pancreatic secretions in vivo change not only from one person to another but also according to the chemical composition of the food, a wide range of physiological relevant flow rates and amylase concentrations were used for these experiments. The flow rate and amylase concentration were varied from 0.3 to 3.0 ml/min and from 2.5 to 25 units, respectively. The levels of these parameters were
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selected based on data from the literature (Mandalari et al., 2010b, Blanquet et al., 2004, Gregg and Sharma, 1978), to simulate slow and fast digestion processes in the small intestine after intake of starch-containing foods. Table 5.4 summarized the experiments performed for starch digestion.

Experiment (I) was performed with the minimum levels of flow and amylase concentration, resulting in low rates of digestion and absorption. In this case, simulated pancreatic secretions (50 ml in total) were injected continuously into the lumen during approximately 2.5 hours at 0.3 ml/min.

On the contrary, experiment (IV) was performed with the highest values selected for both enzyme and flow rate (tenfold higher than those in experiment I). Here, the pancreatic solution was fed into the lumen at an apparent rate of 3.0 ml/min. At this rate, the total pancreatic solution was already injected into the system within the first 20 minutes (16.6 minutes approximately). After that, the concentration of enzyme in the lumen was constant.

Experiments II and III were designed to feed the enzyme at the same ratio either by changing concentration or flow rate, into the lumen side of the SIM. As the purpose of these experiments was to study the effect of amylase on the rate of reaction, the concentration of amyloglucosidase was kept constant (5 U/ml of food) in the pancreatic solution.

5.3.1 Effect of flow rate and enzyme concentration

Digestion was performed in corn starch solutions (1.0%, w/v) previously cooked in boiling water during 30 minutes. Once starch was at room temperature, gastric digestion was
simulated by adjusting the pH of the sample from 5.4 to 2.5 with 1.0 M HCl. After one hour under magnetic agitation, the solution was neutralized with 1.0 M NaHCO₃, increasing the pH from 2.5 to 6.5 to simulate phenomena occurring in the gastro duodenal system. Then, the solution (600 ml) was fed into the lumen side of the SIM. Intestinal digestion started when simulated pancreatic mix (50 ml) was pumped into the lumen side of the SIM and continued for up to 3 hours. Mixing in the small intestine was induced as concentric contractions induced every 2 seconds with the same pattern described in section 4.3. Samples were taken at different time intervals until the end of the experiment to measure reducing sugars, as glucose, by the acid 3,5-dinitrosalicylic acid (DNS) method, in the lumen side of the SIM (inside of the dialysis membrane) and in the recipient side (outside of the membrane). Experiments were done in triplicates from independent experiments as shown in Table 5.4.

Table 5.4 Experimental conditions used for starch digestion.

<table>
<thead>
<tr>
<th>Experiment</th>
<th>Starch (%, w/v)</th>
<th>Enzyme (U/ml)</th>
<th>Flow rate (ml/min)</th>
</tr>
</thead>
<tbody>
<tr>
<td>I</td>
<td>1.0</td>
<td>2.5</td>
<td>0.3</td>
</tr>
<tr>
<td>II</td>
<td>1.0</td>
<td>25.0</td>
<td>3.0</td>
</tr>
<tr>
<td>III</td>
<td>1.0</td>
<td>2.5</td>
<td>0.3</td>
</tr>
<tr>
<td>IV</td>
<td>1.0</td>
<td>25.0</td>
<td>3.0</td>
</tr>
</tbody>
</table>

The results from starch hydrolysis and glucose absorption obtained from the lumen side (membrane) and glucose absorption after generation in the recipient side (outer side) of the SIM varying the concentration of amylase per minute are presented in Figure 5.6 (a) and (b), respectively. An increase in glucose was observed in both sides of the SIM as the concentration of amylase increased over time. Starch hydrolysis exhibited a characteristic “lag phase” in the early stage of the enzymatic reaction followed by a fast rate and decreased rate, reaching steady state after approximately 60 minutes depending on the experimental
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conditions with maximum glucose concentration (~35 mM). Glucose absorption seems to be linear in the recipient side with a maximum of approximately 5 mM after one hour, around ten percent of the glucose released from starch digestion in the lumen. Similar percentages were obtained in experiments of glucose absorption (Chapter 4) for aqueous solutions in which only ten percent of glucose from the lumen moves across the membrane after one hour. Here, this percentage was reached after approximately 120 minutes of digestion, implying that in the first hour, glucose is released from the complex structure of starch (lag phase in Figure 5.6b) and then free sugars with low molecular weight could diffuse to the recipient side of the SIM.

![Figure 5.6 Digestion in the SIM, (a) starch digestion in lumen and (b) glucose absorption in recipient side.](image)

Table 5.5 shows the rate of digestion of starch and glucose absorption calculated from linear regressions of the data plot in Figure 5.6 (a-b). The Overall Mass Transfer coefficients were estimated from the flux of the lumen to the recipient side with the concentration difference at the maximum concentration reached at the steady state in the lumen (31 mM).
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The rate of starch hydrolysis increased up to 90% as the concentration of enzyme increased ten-fold in the lumen side, reaching its maximum value at 25.0 units and 3.0 ml/min while the minimum was observed for the lowest concentration of enzyme at 2.5 units and 0.3 ml/min.

Table 5.5 Digestion and absorption rates for experimental conditions in the SIM.

<table>
<thead>
<tr>
<th>Experiment</th>
<th>Digestion (µM/s)</th>
<th>Absorption (µM/s)</th>
<th>OMTC (µm/s)</th>
</tr>
</thead>
<tbody>
<tr>
<td>2.5 U at 0.3 ml/min</td>
<td>6.91 ± 1.23</td>
<td>0.40 ± 0.04</td>
<td>60</td>
</tr>
<tr>
<td>25.0 U at 0.3 ml/min</td>
<td>9.22 ± 2.34</td>
<td>0.49 ± 0.21</td>
<td>45</td>
</tr>
<tr>
<td>2.5 U at 3.0 ml/min</td>
<td>8.43 ± 0.78</td>
<td>0.51 ± 0.01</td>
<td>45</td>
</tr>
<tr>
<td>25.0 U at 3.0 ml/min</td>
<td>54.55 ± 2.11</td>
<td>0.48 ± 0.05</td>
<td>30</td>
</tr>
</tbody>
</table>

Results in the lumen side also showed no significant difference between experiments II and III, indicating that the method used to feed the enzyme had no significant effect on the rate of starch digestion. This implies that regardless how fast or concentrated the pancreatic solution is fed into the SIM, whether activity of enzyme in digesta is the same the hydrolysis will take place at similar rates of reaction in well-mixed solutions with viscosities like water. Thus, enzyme activity in the lumen can be controlled either by the flow rate at which the pancreatic fluid is fed into the small intestine or the concentration of enzyme present in the digestive secretions.

The increase of enzyme concentration per ml of digesta markedly increased the hydrolysis of starch but had little effect on glucose absorption (Table 5.5). However, the activity of amylase impacted significantly the time at which glucose was absorbed. From the highest enzyme concentration, the change in glucose was detectable at 30 min in the recipient side. There was a ‘delay’ on glucose absorption (from 30 to 60 min) when the rate of starch digestion decreased. Once glucose was generated and released from the starch structure in the
lumen side, absorption took place. Assuming that the chyme moves along the small intestine at 1.0 cm/min (Guyton and Hall, 2006), glucose absorption under conditions presented in Table 5.5, will take place in the first 30, 45 and 60 cm of the duodenum and jejunum. Interestingly, absorption phenomena could be correlated to the point where viscosity of the starch solution in the lumen falls allowing diffusion of nutrients. The glucose profiles in the recipient side appeared to be linear with no appreciable amounts of glucose at the beginning of the experiment.

From the starch digestion experiment it was possible to show the effect that segmentation motion and food formulation had on nutrient delivery to the intestinal wall as a consequence of changes in the mass transfer coefficient. On the other hand, the rate of food digestion might be important in assessing the extension at which they raise blood glucose levels in normal and diabetic people.

5.3.2 Effect of substrate concentration and food formulation

To complement the results above, a series of experiments were performed in the SIM to study the effect that starch concentration has on the rate of glucose absorption. The influence of food formulation was also studied by adding guar gum (0.5 - 0.75%, w/v) to the starch solution. Based on the experimental conditions used in section 5.3.1, preliminary experiments were done to select the ‘optimal’ concentration of amylase and the flow rate at which the enzyme should be injected into the SIM to obtain representative data from the next set of experiments. Table 5.6 summarizes the parameters used.
Table 5.6 Experimental conditions for starch hydrolysis in the SIM

<table>
<thead>
<tr>
<th>Parameters</th>
<th>Experimental values</th>
</tr>
</thead>
<tbody>
<tr>
<td>$x_1$ = Flow rate (mL/min)</td>
<td>1.5 3.0 4.5</td>
</tr>
<tr>
<td>$x_2$ = Enzyme concentration (U/mL)</td>
<td>5 10 15 25</td>
</tr>
<tr>
<td>$x_3$ = Starch concentration (% w/v)</td>
<td>0.5 1.0 2.0 3.0</td>
</tr>
<tr>
<td>$x_4$ = Biopolymer concentration (% w/v)</td>
<td>0.0 0.25 0.50 0.75</td>
</tr>
</tbody>
</table>

These experiments carried out at room temperature and pH 6.5. The levels of the flow rate were selected to represent the range found by Gregg and Sharma, (1975) in human’s pancreatic secretions (4.13 ± 0.88 ml/min). The maximum concentration of amylase (25 U/ml) was selected according to the protocol established by the Institute of Food Research (IFR) in Norwich, UK (Mandalari et al., 2010b).

Digestion was performed at room temperature in 600 ml of digesta consisting of 550 ml of pregelatinized (boiling water/30 minutes) starch solution (1.0%, w/v) and 50 ml of 0.1M Bis-Tris buffer (pH 6.5 ± 0.2). Digestion was monitored in the lumen whereas glucose absorption was monitored in the recipient side of the SIM. Chyme was recirculated along the SIM by a peristaltic pump during 2 hours while the pancreatic solution was fed as shown in Figure 5.1. Mixing of the lumen contents was done by applying segmentation contractions every 2 seconds as described in section 3.2.2. All experiments were done in duplicate or triplicate from independent experiments.

5.3.2.1 Effect of flow rate of the enzyme

In Figures 5.7 (a) and (b) results for starch digestion and glucose absorption at various flow rates are shown. During these experiments the concentration of starch 1.0% (w/v) and the initial concentration of amylase (10 U/ml of food) were constant. Pancreatic solution (50 ml) was fed into the lumen side at 1.5, 3.0 and 4.5 ml/min, meaning that the whole solution (50
ml) was injected after 33, 17 and 11 minutes, respectively. After this period the concentration of the enzymes in the lumen of the SIM was constant.

As expected, the rate of starch digestion increased as the flow rate increased. Starch was hydrolyzed to the extent of 85%, reaching the maximum glucose concentration (31 mM) after 60, 30 and 30 minutes for 1.5, 3.0 and 4.5 ml/min, respectively. Increasing the flow rate from 1.5 to 3.0 has a significant influence in both rate of digestion and time to reach saturation. However when the rate of digestion was increased from 3.0 to 4.5 ml/min, no significant difference was observed neither in rate nor in time of digestion.

In agreement with the results of digestion, no significant variation in glucose absorption was found when the flow rate increased. No glucose was detected in the recipient side during the first twenty minutes of digestion. Interestingly, the delay in glucose corresponds to the lag time in the lumen side. Over this period, glucose increased linearly over time reaching a concentration up to 3.5 mM around 10% of glucose generated in the lumen of the SIM. As no significant difference was observed in digestion and absorption process between 3.0 and 4.5 ml/min in further experiments a flow rate of 3.0 ml/min would be used.
Figure 5.7 Starch digestion and glucose absorption in the (a) lumen and (b) recipient side of the SIM, respectively, varying the flow rate at which the pancreatic solution was fed into the membrane.
5.3.2.2 Effect of the concentration of the enzyme

Pancreatic amylase is present in high activity in the lumen of the small intestine and it has been assumed that the enzyme does not contribute significantly to the rate-limiting process governing carbohydrate digestion and absorption (Slaughter et al., 2001). In order to study the effect of α-amylase concentration upon the reaction rate, a series of experiments were carried out keeping the concentration of starch and the enzyme’s feeding flow rate constant at 1.0% (w/v) and 3 ml/min, respectively. Any change in the amount of product formed over a specified period of time will be dependent upon the level of enzyme present. As Figures 5.8 (a) and (b) show, the rate of digestion and absorption exhibited no significant difference for the range of concentrations of α-amylase used in this set of experiments. The characteristic “lag phase” was observed within the first 10 minutes of the reaction. Prior to this time, the concentration of glucose was less than 2 mM. As time increased the velocity of the reaction increased rapidly reaching a plateau after approximately 30 minutes.

![Figure 5.8](image)

**Figure 5.8** Starch digestion and glucose absorption in the (a) lumen and (b) recipient side of the SIM, respectively, varying the activity of α-amylase.

Samples containing 1.0% (w/v) of starch were digested at room temperature and pH 6.5, whereas pancreatic solution was fed a 3.0 ml/min for all experiments. The velocity of the reaction increased proportionally as the activity of amylase increased. When hydrolysis was done with 5 U/l the rate of digestion was significantly reduced compared with higher
concentrations of enzyme however, no significant differences were found when the enzyme increased from 10 to 25 units (Figure 5.9).

![Graph showing glucose concentration over time for different enzyme concentrations.](image)

**Figure 5.9** Starch digestion in the lumen side of the SIM for various amylase’s activities (5 - 25 U/ml).

Since no significant difference was observed between 15 and 25 units of amylase, in following experiments of starch digestion 15 units per ml of food would be used.

### 5.3.2.3 Effect of concentration of substrate

In Figures 5.10 (a-d) the starch hydrolysis at various initial concentration of starch are presented. Samples were digested at room temperature and pH 6.5, whereas pancreatic solution containing 15 U/ml of α-amylase and 0.5 U/ml of amyloglucosidase was fed at 3.0 ml/min for all experiments. Results showed that the concentration of glucose at enzyme saturation ($C_{inf}$) was influenced by the initial concentration of starch used, increasing as the starch increased from 0.5% to 3.0% (w/v). As the concentration of available substrate molecules increased, concentration of glucose released during amylolysis increased as well.
Chapter 5 - Starch digestion

Over the course of this set of experiments a maximum of 87% of the starch has been converted into glucose (Table 5.7). The percentage of starch transformed can be estimated readily for each initial concentration of starch as follows; whether the molecular weight of the starch monomer \((\text{C}_6\text{H}_{10}\text{O}_5)x\) is 162 g/mol, the maximum concentration for total starch digestion is therefore 19 mM, for 0.5\% (w/v) of starch in 600 ml of digesta. Since the maximum concentration achieved was ~16 mM (Figure 5.10), the percentage of starch digested after 2 hours was approximately 87%.

Table 5.7 Percentage of starch digested after 120 minutes

<table>
<thead>
<tr>
<th>Starch (%,w/v)</th>
<th>Maximum glucose</th>
<th>Percentage of starch digested (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Theoretical (mM)</td>
<td>Experimental (mM)</td>
</tr>
<tr>
<td>----------------</td>
<td>-----------------</td>
<td>----------------------------------</td>
</tr>
<tr>
<td>0.5</td>
<td>18.51</td>
<td>16.20</td>
</tr>
<tr>
<td>1.0</td>
<td>37.04</td>
<td>31.39</td>
</tr>
<tr>
<td>2.0</td>
<td>74.00</td>
<td>56.46</td>
</tr>
<tr>
<td>3.0</td>
<td>111.10</td>
<td>82.27</td>
</tr>
</tbody>
</table>

On average, 80\% of starch digestion was reached under segmentation and in the range of experimental conditions used in the SIM. The differences could be explained due to the increasing viscosity of the sample or inhibition by products. As the concentration of starch increased the viscosity of the solution also increased reducing further interactions between enzymes and substrates. Time course curves in Figure 5.10 suggest that the rate of the reaction was not affected significantly by the initial concentration of starch.
In Figure 5.11 the ratio $C/C_{inf}$ versus time for all concentrations of starch used above was plotted. Interestingly, the curves seem to be very similar regardless the concentration of starch employed in the experiment of digestion. Nevertheless, as the concentration of starch increased there was a delay in reaching equilibrium. For 0.5 – 2.0% (w/v) of starch the maximum concentration at equilibrium was reached at about 30 minutes. However, for 3.0% (w/v) the required time to reach equilibrium was double (60 minutes). This is thought to be
due to the viscosity of the starch solution. Here, diffusion of the enzyme rather than convection took place to reach the surface of the granules of starch.

**Figure 5.11** Starch digestion in the lumen side of the SIM, varying the concentration of starch from 0.5 to 3.0% (w/v).

Results of glucose absorption in the recipient side of the SIM are presented in Figure 5.12. As expected concentration of glucose increased proportionally as the concentration of starch increased resulting in around 10% of the total concentration of sugars generated in the lumen side of the SIM after 2 hours of digestion. In accordance with the results obtained for digestion, at low concentrations of starch, glucose absorption trends were very similar compared to those obtained at high concentration of starch with faster absorption rates (Table 5.8).
Figure 5.12 Glucose absorption in the recipient side of the SIM, varying the concentration of starch from 0.5 to 3.0% (w/v).

Based on data from Figure 5.12, mass transfer coefficients were estimated as described previously. The data were analysed in the linear region of the curve, once the flow was fully developed from 30 to 60 minutes. As the initial concentration of starch increased, glucose in the recipient side and the rate of absorption increased as well. Nevertheless, no significant differences were observed in the mass transfer coefficients for 1.0 – 3.0% (w/v) of starch.

Table 5.8 Rate of absorption and Overall Mass Transfer Coefficient at various starch concentrations.

<table>
<thead>
<tr>
<th>Concentration</th>
<th>Starch (%, w/v)</th>
<th>Glucose saturation (mM)</th>
<th>Absorption ((10^{-7} \text{ M/s}))</th>
<th>OMTC ((10^{-7} \text{ m/s}))</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>0.5</td>
<td>16.20</td>
<td>3.16 ± 0.26</td>
<td>4.19 ± 0.56</td>
</tr>
<tr>
<td></td>
<td>1.0</td>
<td>31.39</td>
<td>5.19 ± 0.22</td>
<td>3.39 ± 0.21</td>
</tr>
<tr>
<td></td>
<td>2.0</td>
<td>56.46</td>
<td>10.08 ± 0.11</td>
<td>3.66 ± 0.19</td>
</tr>
<tr>
<td></td>
<td>3.0</td>
<td>82.27</td>
<td>13.49 ± 0.45</td>
<td>3.36 ± 0.64</td>
</tr>
</tbody>
</table>
5.3.2.4 Effect of viscosity of the digesta

Samples containing 1.0% (w/v) of starch and different concentrations of guar gum (0.25-0.75%, w/v) were digested in the lumen side at room temperature and pH 6.5. Pancreatic solution containing 15U/ml of α-amylase was fed at 3.0 ml/min for all experiments. In Figure 5.13 the effect that the viscosity of the lumen contents had on the rate of glucose absorption is shown. In this particular case, only glucose in the recipient side of the SIM was monitored, because the viscosity in the lumen makes glucose measurement very inaccurate by the colorimetric method employed here. For these experiments, the initial volume of the starch solution in the lumen side was increased from 600 ml in previous experiments to 650 ml due to the viscosity of the solution. The concentration of amylase was adjusted to keep the same concentration of enzyme (15 Units per ml of food).

The rates of glucose absorption were significantly reduced as a result of increasing the viscosity of the lumen contents by adding guar gum. When initial viscosity was increased from 0.3 (0.5% w/v guar) to 7.0 Pa.s (0.75% w/v guar) glucose concentration was reduced up to 45%. Moreover, compared with results in Figure 5.12 for 1.0% (w/v), glucose absorption was delayed. Glucose could be monitored in the recipient side after 20 minute of starting digestion (Figure 5.12), however, for viscous solutions glucose was measured after 30 minutes the reaction started (Figure 5.13). As viscosity of the digesta increased interactions between substrates and enzymes were impaired, resulting in a reduction of glucose released from starch structure.
Figure 5.13 Glucose absorption in the recipient side of the SIM, varying the concentration of guar gum from 0.25 to 0.75% (w/v).

The rate of glucose absorption and mass transfer coefficients are shown in Table 5.9. Both were reduced as the viscosity of digesta increased. Restrictions in mass transfer due to variations of viscosity cause a reduction in the rates of diffusion of substrate and products associated with digestion processes in the small intestine.

These results give insights about physical and chemical phenomena occurring in the small intestine and also how rates of digestion and absorption are impacted, based on food formulation and fluid properties of the chyme. Again, these trends are consistent with the *in vivo* findings that viscous fibres reduced postprandial glucose levels (Jenkins et al., 1982).

A comparison between the OMTC obtained for starch-guar solutions and those calculated for glucose-guar solutions in Chapter 4 (section 4.3.2.1) is presented in Table 5.9.
Table 5.9 Rate of absorption and Overall Mass Transfer Coefficient at various concentrations of guar. Concentration in the lumen side at enzyme saturation was 31.35 mM.

<table>
<thead>
<tr>
<th>Guar concentration (%)</th>
<th>Absorption (10^{-7} M/s)</th>
<th>OMTC starch (10^{-7} m/s)</th>
<th>*OMTC glucose (10^{-7} m/s)</th>
</tr>
</thead>
<tbody>
<tr>
<td>0.00</td>
<td>5.84 ± 0.12</td>
<td>3.82 ± 0.33</td>
<td>4.56 ± 0.02</td>
</tr>
<tr>
<td>0.25</td>
<td>5.42 ± 0.52</td>
<td>3.54 ± 0.16</td>
<td>4.66 ± 0.39</td>
</tr>
<tr>
<td>0.50</td>
<td>3.74 ± 0.17</td>
<td>2.73 ± 0.19</td>
<td>3.47 ± 0.05</td>
</tr>
<tr>
<td>0.75</td>
<td>3.26 ± 0.34</td>
<td>2.13 ± 0.04</td>
<td>3.43 ± 0.01</td>
</tr>
</tbody>
</table>

*Results taken from section 4.3.2.1 from glucose solutions added with guar gum at different concentrations.

As results show, increasing viscosity of the starch solution reduced mass transfer by up to 45%. However, when glucose-guar solutions were evaluated in Chapter 4, absorption was only reduced 25%. The OMCT for glucose-guar solutions were higher than those obtained after digestion of starch added with similar concentrations of guar gum. Compare to glucose solutions in which glucose is already available, adding soluble fibres to starch had a significant impact on glucose absorption. In starch solutions, glucose needs to be released first from the complex matrix of starch to be able to cross the membrane then. However, the enzyme has an additional physical barrier formed by guar.

These results suggest that adding viscous polymers to foods inhibit not only the absorption of glucose from the intestinal lumen to the wall but also to diffusion of enzyme to reach starch molecules.
5.4 Michaelis – Menten kinetics

Michaelis-Menten equation was used to characterise the behaviour of reaction in which the enzyme concentration is remarkably small compared to the substrate concentration (Dona et al, 2010). The main principle of Michaelis – Menten reaction is as follow:

\[ E + S \leftrightarrow ES \rightarrow E + P \]

where \( E \) is the concentration of the enzyme, \( S \) refers to the substrate and \( P \) refers to the product.

The Michaelis-Menten equation in the differential form can be used to describe the dynamics of substrate depletion as

\[
\frac{dS}{dt} = \frac{-V_{\text{max}}S}{K_m + S} \tag{5.3}
\]

where \( S \) is the substrate concentration, and \( V_{\text{max}} \) and \( K_m \) are the maximal rate and Michaelis half saturation constant, respectively. Equation 5.3 can be readily integrated to obtain the integral form of the Michaelis-Menten equation

\[
K_m \ln\left(\frac{S_o}{S}\right) + S_o - S = V_{\text{max}} t \tag{5.4}
\]

where \( S_o \) is the initial substrate concentration. Equation 5.4 is not linear and clearly implicit with respect the substrate concentration.

Numerical approaches such as bisection and Newton-Raphson methods are necessary to calculate \( S \). In order to obtain the explicit form of Equation 5.4, Goudar et al. (2004) rearrange to form:

\[
S + K_m \ln(S) = S_o + K_m \ln(S_o) - V_{\text{max}} t \tag{5.5}
\]

Substituting \( \phi = S/K_m \) in Equation 5.5 results in
\[ \phi K_m + K_m \ln(\phi K_m) = S_o + K_m \ln(S_o) - V_{\text{max}} t \]  

(5.6)

Dividing Equation 5.6 by \( K_m \) and rearranging results in

\[ \phi + \ln(\phi) = \frac{S_o}{K_m} + \ln\left(\frac{S_o}{K_m}\right) - \frac{V_{\text{max}} t}{K_m} \]  

(5.7)

The left hand side of Equation 5.7 is analogous to the Lambert \( W \) function (Corless et al., 1996).

\[ W(x) + \ln\{W(x)\} = \ln(x) \]  

(5.8)

where \( W \) is the Lambert \( W \) function and \( x \) the argument of \( W \) from equations and expression for \( \phi \) may be obtained as

\[ \phi = W\left\{\frac{S_o}{K_m} \exp\left(\frac{S_o-V_{\text{max}} t}{K_m}\right)\right\} \]  

(5.9)

\[ S = K_m W\left\{\frac{S_o}{K_m} \exp\left(\frac{S_o-V_{\text{max}} t}{K_m}\right)\right\} \]  

(5.10)

Linearization

\[ \frac{t}{\ln(S/S_o)} = \frac{1}{V_{\text{max}}} \frac{(S_o-S)}{\ln(S/S_o)} + \frac{K_m}{V_{\text{max}}} \]  

(5.11)

\[ \frac{(S_o-S)}{\ln(S/S_o)} = V_{\text{max}} - \frac{K_m}{t} \ln(S/S_o) \]  

(5.12)

\[ \frac{t}{(S_o/S)} = \frac{K_m}{V_{\text{max}}} \frac{\ln(S/S_o)}{(S_o-S)} + \frac{1}{V_{\text{max}}} \]  

(5.13)

This approach involves the use of substrate depletion rather than initial velocity-substrate concentration.

From Equations 5.11 to 5.13 it was possible to estimate the initial values of the kinetics parameters \( V_{\text{max}} \) and \( K_m \), from the progress curve analysis. These parameters were used to
calculate the resulting in the curve of Figure 5.13. Fit was not very good by this method, needing to look into more detail to compare with kinetic parameters reported in literature.

![Graph](image_url)

**Figure 5.14** Simulated substrate depletion data (cross) along with the model predictions (solid line) for 1.0% (w/v) starch digestion in the STR at concentrations of enzyme constant.

\[
V_{\text{max}} = 0.59 \text{ mM/min}
\]

\[
K_{\text{m}} = 9.02 \text{ mM}
\]

5.5 Conclusions

This research has demonstrated the capability of using an *in vitro* rig to simulate and obtain an engineering understanding of transport phenomena occurring in the small intestine leading to absorption of active components.

By using the STR, starch digestion has a fast rate of reaction (44.8 µM/s). However, the hydrolysis of starch in the SIM has a sigmoidal behavior with a rapid rate (24 µM/s) after the first ten minutes of digestion and with a ‘lag phase’ at the beginning of the reaction due to the
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incomplete mixing of the starch and amylase at the early stage of the hydrolysis within the first ten minutes.

The effect of the presence of guar showed a 36% reduction on the mass transfer coefficient when mixing or segmentation was applied. Overall, the rate of enzymatic reactions was reduced when guar (0.5%, w/v) was added to starch solutions and bread, delaying the access of actives to the absorptive epithelium due to a decrease of both propulsion and mixing.

The rate of starch hydrolysis increased up to 90% as the concentration of enzyme increased ten-fold in the lumen side, reaching its maximum value at 25.0 units and 3.0 ml/min while the minimum was observed for the lowest concentration of enzyme at 2.5 units and 0.3 ml/min. The velocity of the reaction increased proportionally as the activity of amylase increased. As the concentration of starch increases the viscosity of the solution also increases. In this case, for high concentrations of starch (3.0%, w/v), diffusion of the enzyme rather than convection took place to reach the surface of the granules of starch.

Overall, the rate of glucose absorption and mass transfer coefficients were reduced as the viscosity of digesta increased. Restrictions in mass transfer due to variations of viscosity cause a reduction in the rates of diffusion of substrate and products associated with digestion processes in the small intestine.

From the starch digestion experiments present in this Chapter it was possible to show the effect that segmentation motion and food formulation had on nutrient delivery to the intestinal wall as a consequence of changes in the mass transfer coefficient. On the other hand, the rate of food digestion might be important in assessing the extension at which they raise the blood glucose levels in normal and diabetic people.
CHAPTER 6 - MIXING PROCESSES IN THE SIM

6.1 Introduction

Food to be processed in the gastrointestinal tract requires an adequate degree of mixing. Particularly, in the small intestine, mixing is a result of segmentation contractions that keep solid food particles dispersed in the chyme and promote chemical reactions between substrates and enzymes. Mixing is responsible for mass transfer enhancement and is crucial for absorption. As established in Chapter 4, mixing or segmentation increased glucose absorption up to 30%. Isolated longitudinal contractions enhance mass transfer processes in the small intestine by providing radial mixing of the chyme (Schulze and Clark, 2008, Macagno et al., 1982). Most studies of flow motion in the small intestine point out that oscillatory flow promotes ‘turbulent mixing’, in which radial velocity components are significant (Mackley and Stonestreet, 1995). Long stroke oscillations, for example, have shown to improve heat and mass transfer as a result of fluid mixing (Howes et al., 1991).

In the gastrointestinal tract, various types of movements have been identified. Retrograde jets and oscillatory flow vortices, for example, mix two separate fluid boluses enhancing mass transfer in the lumen. As few experimental preparations exist to record the complex flow that underlie digestion and absorption processes (Schulze and Clark, 2008) in the small intestine, this Chapter describes visual flow patterns occurring as a result of mixing or segmentation contractions in the SIM. Flow visualization techniques such as PLIF (Particle Laser-Induced Fluorescence) and PEPT (Positron Emission Particle Tracking) were used to identify fluid patterns and velocities profiles of Newtonian and non-Newtonian model fluids in the SIM.
6.2 Optical flow visualization techniques

6.2.1 Preliminary flow visualization experiments

To study flow patterns in the SIM as a function of segmentation contractions, a series of preliminary experiments were carried out in water and CMC solutions (1.0%, w/v). In order to visualize the motion of these fluids, a red food coloring was injected into the lumen side of the SIM. The movements of the dye were recording for about 1 - 2 minutes by a high definition dual camera (Sanyo, Xacti HD VPV-FH1). The camera was located in front of the SIM, between the two segmentation cuffs area. For these experiments, the dialysis membrane was substituted by a flexible transparent polyethylene tube with the same diameter and length of the membrane used in previous experiments: 0.03 m. Representative frames were extracted at selected times from each video to study the flow patterns of the dye. After filling the lumen side with the test solution, either water (control) or CMC (1.0%, w/v), the red dye was fed at a flow rate of 3.0 ml/min into the centre of the intestinal segment by means of a tubing (0.003m of diameter) simulating the pancreatic duct as shown in Figure 6.1(a). The rig used in this section to feed the dye was similar to that used in Chapter 5 (Figure 5.1) to feed the enzyme solution into the lumen side of the SIM. The main purpose of these experiments was to track the path that amylase (represented by the red dye) follows during digestion as a function of both segmentation and food formulation. Fluid motion was studied at experimental conditions listed in table 6.1.

Table 6.1 Experimental conditions for studying flow patterns of the dye in the SIM

<table>
<thead>
<tr>
<th>Fluid</th>
<th>Experiment I</th>
<th>Experiment II</th>
<th>Experiment III</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Newtonian:</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Water</td>
<td>Steady state</td>
<td>Net flow induced by a peristaltic pump</td>
<td>Segmentation induced by inflatable cuffs</td>
</tr>
<tr>
<td><strong>Non Newtonian:</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>CMC (1.0%, w/v)</td>
<td>Steady state</td>
<td>Net flow induced by a peristaltic pump</td>
<td>Segmentation induced by inflatable cuffs</td>
</tr>
</tbody>
</table>
Segmentation contractions were generated at a rate of ten cycles per minute (selected for being physiological relevant in the small intestine (Guyton and Hall, 2006)) with the following standard pattern: 2 seconds inflation, 2 seconds deflation and 2 seconds delay between each cuff. As previously described in section 3.2, two cuffs located in the proximal and distal ends of the intestinal segment (50 cm of length) were used to generate sequential contractions along the SIM.

As shown in Figure 6.1 (a) when the dye was injected in the SIM in the absence of bulk flow and segmentation (experiment I), the dye exhibits spontaneous diffusion through water. When net flow of the fluid was generated by a peristaltic pump (3 rpm), the diffusion of the dye into water was accelerated, displaying segregation and a characteristic parabolic flow pattern in which the maximum velocity was observed, as expected, in the centre of the pipe (radius equal to zero) (Figure 6.1b).

Overall, reverse flow in the SIM occurs both during the contraction and during the relaxation of the segment and as a result, the lumen content is constantly refilling fluid from the reservoir. Similarly, reverse flow and refilling of cleared segments are regularly observed in the small intestine in situ (Schulze and Clark, 2008).

**Figure 6.1** Diffusion of the dye through the SIM in water; (a) steady state and (b) propulsion by a peristaltic pump (volumetric flow of $1.2 \times 10^{-5} \text{ m}^3/\text{s}$), after one minute of the dye injection.
Chapter 6. Mixing processes

As segmentation progresses, the dye is moving forward and backward within the fluid promoting striation of the dye. Once the contraction stops (relaxation of the lumen) the dye returns to the initial point of injection. During relaxation, convective mixing stops and diffusion of the dye occurs until the next contraction. These periodical movements keep mixing the dye throughout the fluid. The cycle of contractions distributes and disperses the dye through the intestinal segment in which simultaneous reduction of intensity and scale of segregation occurs over time.

In Newtonian fluids such as water, segmentation contractions generate strong intraluminal flow, inducing retropropulsive and recirculating (oscillatory) flows that contribute to the mixing processes in the small intestine. Whilst backflow helps to mix the dye over the long axis of the SIM, flow vortices or eddies distribute the dye (enzyme and food particles) over the radius of the SIM (Figure 6.2).

![Figure 6.2](image-url) Mixing flow patterns of the dye in the SIM when segmentation occurs in water. The direction of the flow imposed by each contraction is represented by the block arrows.
Chapter 6. Mixing processes

Although both contribute to mass transfer phenomena, it is though that the latter contributes the most by promoting radial mixing of the lumen contents. Retrograde and radial flows promote mixing and absorption of active molecules in the small intestine by generating turbulent mixing in the lumen, helping to distribute food particles over the surface of the intestine and promoting enzyme-substrate interactions.

Segmentation contractions produced turbulent mixing, creating vortices along the radial axis of the flexible tube. The occurrence of flow vortices was seen during experimentation and that further evidence on this is provided later when PEPT results were presented in section 6.2.2. Vortices promote radial mixing reducing the mixing time required to obtain a homogeneous solution along the length of the tube (axial and radial axis). These flow events are essential during digestion and absorption processes increasing mass transfer in the gut. For instance, when solid particles sediment in the small intestine due to their particle size, contractions keep food particles of the chyme dispersed, helping to interact with intestinal and pancreatic secretions improving mass transfer in the intestine. Similar flow events were already studied in the duodenum and ileum of guinea pigs in the research conducted by Schulze-Delrieu (2008).

For CMC solution (1.0%, w/v), minimal dye dispersion (Figure 6.3a) was observed as a result of increasing the viscosity of the fluid. Although diffusion was increased when a peristaltic pump was used (Figure 6.3b), the dispersion of the dye was not as fast as it was for water due to the increasing viscosity.
Experiments for 1.0% (w/v) CMC solutions in which segmentation was applied are shown in Figure 6.4. Again the dye moves back-and-forth accordingly to the contraction applied to the SIM. Although segmentation seems to bring dye in the radial direction once contractions occlude the lumen side of the SIM, no mixing occurred when the contraction vanished and the viscosity (zero shear viscosity) of the bulk solution increased. As segmental contractions are isolated local contractions no permanent mixing is induced in the lumen for laminar flow. Unlike water, when the viscosity of the chyme increased the dye moves as a compact volume. Once the contraction stops the dye remains in the middle of the SIM static, with no further movement until the next contraction was generated. Although successive contractions tend to mix more and more, clearly viscous solutions require more time to mix and create homogeneous solutions.
Figure 6.4 Mixing flow patterns of the dye in the SIM when segmentation occurs in 1.0% (w/v) CMC. The direction of the flow imposed by each segmentation contraction is represented by the block arrows.

As segmentation progresses the intensity of mixing increases. After one minute, intensity and scale of segregation was higher in CMC solutions than segregation in water as Figure 6.2 revealed. For water, for example, the distribution of the dye was homogenous after one minute, on the contrary, when the viscosity of the solution increases mixing takes longer. In Figure 6.5, a comparison of the distribution of the dye into water and CMC solutions (0.5 and 1.0%, w/v) after one minute was presented. Clearly the distribution of the dye was more homogeneous in water than in CMC solutions. This is thought to be caused by a reduction of convective mixing in viscous fluids. Viscous solutions resist the effect of gastro-intestinal contractions, inhibiting both propulsion and mixing (Read and Welch, 1985).
Figure 6.5 Dispersion of the dye in (a) water and (b) CMC (1.0%, w/v) after one minute of applied segmentation in the SIM at 2s.

This behaviour could aid the explanation of the results found for glucose absorption in Chapter 4 and 5, in which, absorption rates and mass transfer coefficients were higher for aqueous solutions compare with those for guar and CMC. Furthermore, when segmentation took place in the SIM, glucose absorption was increased up to 30% for water. On the contrary, when the viscosity of the solution increased absorption increased as well, but only 10% under segmentation.

Overall, by increasing the viscosity of the chyme is possible to reduce interactions between enzyme-substrate and intestinal secretions; (i) reducing the rate of digestion and (ii) delaying the access of actives molecules to the epithelium.

6.2.2 Planar Laser Induced Fluorescence (PLIF)

PLIF is a visualization flow technique that gives qualitative and quantitative information on the mixing effectiveness. In this technique the dispersion of a fluorescent dye into the bulk fluid is based on the area of the tracer immersed in the solution as a result of a mixing mechanism. To complement the above results, PLIF was used to determine the effect of mixing of 0.5 ml of Rhodamine dye into viscous CMC solutions (0.5 and 1.0%, w/v) as a result of the segmentation contractions applied to the SIM. As PLIF technique requires a transparent system the original geometry of the SIM was modified by introducing a Perspex
flow visualization cell of 20 cm long and 5 cm both deep and wide (section 3.5). Experiments were carried out as before using the characteristic two cuffs set up with 2s inflation, 2s deflation and 2s delay times. The air pressure used for the inflation was 0.5 barg. The net flow rate was set at approximately 1.0 cm per minute, as established by Guyton and Hall (2006) for the small intestine. This was achieved using a peristaltic pump to give a volumetric flow of $1.2 \times 10^{-5} \text{m}^3/\text{min}$ (velocity of 1.6 cm/min), which was the minimum flow rate of the pump. For these experiments, the injection of the fluorescence dye was done manually by means of a 1.0 ml syringe and needle. The needle was previously fixed into the visualization cell to inject the Rhodamine at the centre of the Perspex tube to avoid diffusion of the dye before segmentation contractions start. Since produced clear solutions, CMC was selected in PLIF flow visualization experiments. Experiments were done in water and CMC solutions (0.5 and 1.0%, w/v) at two different mixing frequencies (one and two seconds). The images taken were in the form of a digital picture (as seen in Figure 6.11 (a) and all the images for each experiment were taken from time zero (when full injection of the dye had taken place) until approximately 60 seconds, at an image acquisition rate of 2 Hz. This provided 80 images that were analysed for each experiment using MATLAB.

Figure 6.6 presents a typical sequence of images that were recorded for a 0.5% (w/v) CMC solution after the first contraction was fully developed. The clear circular area in each image represents the Rhodamine dye injected into the SIM. The purpose of these experiments was to model the movement of a discrete bolus (represented by the dye) in the centre of the lumen as a function of both mixing and viscosity of the fluid. As contraction develops the dye moves within the fluid from the centre of the pipe to the distal part of the segment (Figure 6.6). As the first contraction vanishes the lumen segment relaxes and the dye returns at the initial position of injection (back and forth movement).
Figure 6.6 Dye bolus motion as a left contraction develops in the SIM, images captured by Planar Laser Induced Fluorescence (PLIF) for 0.5% (w/v) CMC.
When the cuff in the left occluded the flexible wall of the SIM, fluid moves faster through the length of the SIM and as a consequence the dye starts to mix due to motion. As the cuff deflates the dye returns slowly to the original location in the centre of the pipe. After approximately 4 seconds, the second cuff in the right starts to inflate and the particle moves back along with the volume of fluid displaced. These periodic contractions help not only to mix the chyme along the SIM but also to increase the residence time of the food, maximizing

Figure 6.7 displays three representative images of the dye movement in 0.5% (w/v) CMC after the first set of segmentation contractions take place. The tiny bolus simulated by the clear circle (2D) in the middle of the picture moves according to the left or the right segmentation contraction. Images show the characteristic retropulsive or pendular movements found in the small intestine by segmentation contractions (Macagno and Chistensen (1982)). Once segmentation started, the bolus moves accordingly back and forward. The displacement of the dye is a function of the resistance of the fluid to movement (viscosity). In the control solution (water) the dye was quickly dispersed with a reduction of the area of the initial sphere shape (data are not shown). As segmentation continues it was possible to identify three characteristics zones:

i. **Frames in the left.** Maximum displacement after first contraction occurs from the centre to the right, in the same direction as contraction occurs (Figure 6.7 (c) and (f)).

ii. **Frames in the middle.** Relaxation of the lumen no segmentation was applied to the SIM (Figure 6.7 (b) and (e)).

iii. **Frames in the right.** Maximum displacement after second contraction occurs from the centre to the left, opposite direction of the contraction (Figure 6.7 (a) and (d)).
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Figure 6.7 (a) and (b) shows that the displacement of the dye is higher when the duration of the contraction increases from 1 to 2 seconds. Whilst the dye moves from 0.5 to 0.6 along the length of pipe when contractions last only one second, the displacement increases from 0.5 to 0.7 when the contraction last 2 seconds. As frequency increases the volume displaced by the cuff increases and the contraction is shallow. This is in accordance with the results found in section 4.3.3 in which mass transfer coefficient were higher when the mixing frequency was increased from 1 to 2 seconds in CMC solutions.

At time zero the dye was injected, after 2 seconds of delay the contraction in the left develops and after 4s the dye reaches the maximum displacement in 0.7 (Figure 6.7 c,d). As the contractions vanishes (6s) the dye returns to the centre of the pipe and remains in the centre during 4 more seconds (t = 10s) in this position until the next contraction (right cuff) develops t =10 s and it is hold for 2s (t =12 s) 0.2 more and the cycle starts again.

Figure 6.7 Dye bolus motion for a set of segmentation contractions for 0.5% (w/v) CMC solution; contractions generated every one (a-c) or two seconds (d-f).
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Images also defined how the contraction stretched the gray bolus within the surrounding black fluid, separated the bolus into portions, and replaced the original small interface between bolus and fluid with multiple large interfaces of layers.

For CMC 1.0% (w/v), however, deformation of the dye was not as significant as it was for 0.5% (w/v) CMC solutions. Again when the mixing frequency increased the displacement was increased (Figure 6.8). Overall the first cuff was not as effective in mixing as the second cuff by the action of the peristaltic pump which limited the back flow. As it can be seen in Figure 6.8c, the segmentation contraction in the left produced a major displacement, than the contraction in the right because in this particular case, the flow motion comes in the same direction of the flow. Thus, when the right cuff is inflated, the movement of the dye bolus from right to left but with a minor displacement, because in this case the movement is against the net flow as previously said. As such both propulsion and mixing contribute to the bulk flow bringing particles of food and enzymes together and creating a better distribution of the dye in the fluid.

![Figure 6.8](image)

Figure 6.8 Dye bolus motion for a set of segmentation contractions for 1.0% (w/v) CMC solution; contractions generated every one (a-c) or two seconds (d-f).
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From these images it is evident that the viscosity of the fluid and the frequency of contractions affect the effectiveness of mixing and as a consequence mass transfer phenomena in the small intestine.

6.3 Positron Emission Particle Tracking (PEPT)

The PEPT technique is a powerful non-intrusive 2D and 3D flow visualisation tool that allows tracking a labelled tracer particle in opaque multiphase systems that cannot be studied by optical methods. With PEPT, the tracer is located by means of a dual positron camera and an algorithm for computing the tracer location (Barigou, 2004, Fangary et al., 2000, Parker et al., 2002).

PEPT is a variation of the Positron Emission Tomography (PET) used in medical studies to evaluate physiological functions of the body by introducing a positron emitting radioactive tracer (Barigou, 2004). PEPT has been used successfully to study mixing patterns in powders, semi-solids and liquids (Bakalis et al., 2004). This technique is particularly useful for the study of multiphase flows, to map the flow of fluids and particles (Barigou, 2004).

To monitor the flow through the small intestine a single label tracer (600 µm) was injected into the SIM as shown in Figure 6.9. These experiments were carried out in the opened SIM configuration in which fluid was re-circulated by a peristaltic pump at 1.2 x 10⁻⁵ m³/s (see section 3.5.2). As in section 6.2, the dialysis membrane of the system was substituted with a polyethylene flexible tube of 0.03 m of diameter and 0.40 m of length.

Although the equipment and the camera were aligned the axis of the camera does not necessarily coincide with the coordinate system of the SIM.
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Figure 6.9 Schematic of the experimental set up used for PEPT.

Three different conditions were studied, the effect of: viscosity of the biopolymer (guar gum 0.5 and 1.0%, w/v), frequency of segmentation (1s and 2s) and the net flow rate (0 and 3 rpm). Experiments were done at room temperature and 0.5 barg inlet air pressure for each cuff. Table 6.2 summarizes the experimental conditions used for PEPT. During experiments the activity of the tracer was monitored to ensure proper detection. In this case, no control solution was used because the activity of the tracer decays in aqueous solutions.

Table 6.2 Experimental conditions used for PEPT

<table>
<thead>
<tr>
<th>Assay</th>
<th>Guar gum concentration (%, w/v)</th>
<th>Net flow (ml/min)</th>
<th>Segmentation contractions</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>0.5</td>
<td>No net flow</td>
<td>2s</td>
</tr>
<tr>
<td>2</td>
<td>0.5</td>
<td>12.0</td>
<td>2s</td>
</tr>
<tr>
<td>3</td>
<td>0.5</td>
<td>12.0</td>
<td>1s</td>
</tr>
<tr>
<td>4</td>
<td>1.0</td>
<td>No net flow</td>
<td>2s</td>
</tr>
<tr>
<td>5</td>
<td>1.0</td>
<td>12.0</td>
<td>2s</td>
</tr>
<tr>
<td>6</td>
<td>1.0</td>
<td>12.0</td>
<td>1s</td>
</tr>
</tbody>
</table>
Chapter 6. Mixing processes

The experimental flow conditions are explained below:

- No net flow induced by peristaltic pump and local flow from segmentation contractions by cuffs ‘1’ and ‘2’ alternatively with a two second inflation time, two second deflation time and two second delay.
- Net flow induced by a peristaltic pump and local flow from segmentation contractions by cuffs ‘1’ and ‘2’ alternatively with a two second inflation time, two second deflation time, two second delay, generating 10 contractions per minute (0.166 s\(^{-1}\)).
- Net flow induced by a peristaltic pump and local flow from segmentation contractions by cuffs ‘1’ and ‘2’ alternatively with one second inflation time, one second deflation time, one second delay, generating 20 contractions per minute (0.333 s\(^{-1}\)).

Both set of contractions, 10 and 20 contractions/minute, were chosen in a physiological context in which contractions occurring in the duodenum are simulated in (i) normal subjects (10/min) according to those reported by (Guyton and Hall, 2006) and (ii) when gastrointestinal disorders such as gastroenteritis or intestinal obstruction affect the bowel motility, producing weak contractions and increasing in some cases the contractions observed in the duodenum up to 20/min.

PEPT produces a set of particle trajectories showing how tracer moved thought the SIM as a consequence of the segmentation contractions. PEPT data include locations of the particle in ‘x’, ‘y’ and ‘z’ coordinates as a function of time. In the SIM the axial (v\(_x\)) and radial velocity (v\(_r\)) components are of importance on mixing of the lumen content. The velocity in the SIM is a contribution of both segmentation (local velocity) and the net flow displacement produced by the peristaltic pump (net flow velocity) (Equation 6.1).
Chapter 6. Mixing processes

\[ v = v_{osc} + v_{ave} \]  

(6.1)

where \( v_{osc} \) is the oscillatory velocity induced by the segmentation contractions in the SIM and \( v_{ave} \) is the average velocity due to segmentation and peristalsis (net flow velocity).

The velocity component along the axis of the SIM \( (v_x) \) is related to the particle path by Equation (6.2)

\[ v_x = \frac{\Delta x}{\Delta t} = \frac{x_2-x_1}{t_2-t_1} \]  

(6.2)

where \( x \) is the axial coordinate and \( t \) is time.

Using regression analysis a line was fitted to a number of the ‘x’ locations of the particle versus time. The slope of the line was the axial velocity component of the particle.

Figures 6.10 (a) and (b) show a characteristic retropulsive flow pattern in the ‘x’ direction of the tracer for guar solutions (0.5%, w/v), as a result of the segmentation contractions generated by the second cuff of the SIM (see Figure 6.9). In Figure 6.10 (a), no net flow was induced by the peristaltic pump, i.e. the motion of the particle was induced only by segmentation. Since the particle was located in the last cuff of the system, the effect of the first cuff was not observed in this experiment (flat part of the graph). Figure 6.10 (a) shows that the tracer remains essentially in the same cross channel position. On the other hand, when the net flow of the fluid was induced by the peristaltic pump (Figure 6.10b) the particle moved little faster along the ‘x’ axis. In both cases the back-and forth movement of the particle was clearly observed due to the contraction produced by distal cuff of the SIM. Particle moves within an average velocity in the x-axis at 2.0 cm per minute due to the contribution of both the net and the local flows induced by the peristaltic pump and the segmentation contractions, respectively. This value is in accordance with those reported by
Chapter 6. Mixing processes

Guyton and Hall (2006) in the human small intestine. Although little movement was induced by segmentation this is essential to mixing processes taking place in the intestine.

![Figure 6.10](image_url)

**Figure 6.10** Flow path of a radioactive tracer (600 μm) under segmentation contractions in the SIM for 0.5% (w/v) guar solution (a) segmentation and (b) segmentation and peristalsis.

As Figure 6.11 shows, it is possible to follow the trajectory of the radioactive tracer, in which three distinctive zones were identified according to the position of the particle:

**Zone A. The tracer is placed at the end of the first cuff of the SIM (50-100s)**

As first contraction occurs (cuff 1), the particle moves in the positive “x” direction with a displacement from ~75 to 150 mm for about two seconds (duration of the contraction). Once deflation of cuff 1 starts, the particle moves backward and returns to its initial position. As the particle is distant from the second cuff the movement of the particle seems to be unaffected by the cuff on the right (cuff 2), i.e. a very small displacement in the negative direction of the “x” axes is shown, represented as a small down peak.
Zone B. Tracer is located in the centre of the SIM (150-250s)

At this point the particle has moved through the length of the SIM and it is placed in the middle of the SIM. This is a result of both peristalsis and segmentation motion. In this case, the effect of both cuffs appeared represented by the two characteristic peaks that are approximately at the same order of magnitude. As the particle moves forward over time along the length of the SIM, the down flow peak is fully developed and begins to increase. On the contrary, the peak related to cuff 1 starts to vanish indicating that the particle is closer to the second cuff and its back-and-forth movement is only dependable on the second cuff. As experiment progressed over time, the effect of the first contraction was not significant any longer as shown in the schematic below (Zone C in Figure 6.11).

Zone C. Tracer is near to the second cuff of the SIM

Once the particle is closed to the second cuff, the behavior observed in zone “A” is inversed and the particle moves as a result of contractions produced by the second cuff. For that reason the peak in the negative direction of “x” increased in magnitude compare to the peak produced by the first cuff (in the left) which was reduced. The movement of the particle is significantly affected by the second cuff while the first cuff does not have an effect on the movement of the particle.

Therefore, the trajectory of the particle depends on its location. As described before, the flow pattern of the tracer was slightly different when the particle was located close to the first cuff of the SIM for 0.5% (w/v) guar solutions. Interestingly, in this experiment the tracer was following the dynamics of the two contractions, produced by cuffs 1 and 2, and the net flow induced by the pump.
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First contraction    Second contraction

Figure 6.11 Flow path of the tracer over time under segmentation contractions generated every 2 seconds in the SIM for 0.5% (w/v) guar solution. Schematic ‘A-C’ represents the relative position of the particle as a function of both cuffs.

The tracer patterns show that the strongest fluid motions are produced under segmentation with velocities of oscillation up to 0.05 m/s (Figure 6.11). This experimental measured velocity is in agreement with the theoretical value (0.06 m/s), estimated in Chapter 4.

The motion of the fluid induced by segmentation contraction waves generates recirculating vortices or eddies that promotes turbulence, mixing the lumen content. To show clearly the
recirculating flow pattern due to a segmentation contraction, a 3D plot was built for time, x-axis and y-axis data from isolated contractions in Figure 6.12. The x-axis and y-axis represent the axial and the radial position of the tracer, respectively, whereas the time (in seconds) is represented by the color bar aside in Figure 6.12.

![Figure 6.12](image_url)

**Figure 6.12** Characteristic eddy identified in the SIM as a result of a segmentation contraction in guar gum (0.5%, w/v).

As shown in Figure 6.12, oscillatory flow induced by segmentation, is an efficient way to mix the lumen contents along the radial y-axis of the SIM, transporting particles laterally towards the intestinal wall. As the contraction develops (A-B) from 92 to 94 seconds (2s duration), the fluid moves rapidly along the y-axis ~ 1.0 cm of the diameter of the tube from position A to B, once the contraction vanishes, the particle returns to its initial position (B-C).
Phenomena observed during experimentation suggest that segmentation movements are responsible for oscillatory flow that brings dissolved food material from the centre of the small intestine to the wall (A-B). Thus, segmentation affects the rates at which digestion and absorption take place in the intestine, it was also demonstrated that mass transfer processes are enhanced by segmentation motion.

As Figure 6.13 shown, four consecutive peaks were extracted from Figure 6.11 to identify formation of vortices or eddies generated by oscillatory flow in the SIM when segmentation was applied.

Figure 6.13 Flow path induced under segmentation in guar gum solutions (0.5%, w/v).

Results are displayed in Figures 6.14 and 6.15 for each peak at (a) 56, (b) 67, (c) 80 and 92 seconds. In Figure 6.16 all of these recirculating eddies are shown together (identified by colour).
Figure 6.14 Characteristic eddies identify in guar gum solutions generated as a result of the segmentation contractions (a) 56-60 seconds and (b) 67-71 seconds.
Figure 6.15 Characteristic eddies identify in guar gum solutions generated as a result of the segmentation contractions (a) 80-84 seconds and (b) 56-84 seconds.
Chapter 6. Mixing processes

Figure 6.16 Characteristic eddies identify in guar gum solutions (0.5%, w/v) generated as a result of segmentation contractions (a) 80-84 seconds and (b) 56-84 seconds.

Segmental flow produced vortex formation cycles (oscillation) (Figures 6.14-16) which enhanced mixing and dispersion of chyme in the SIM. As a result, mass transfer processes are enhanced in low-viscosity fluids. This could explain why the OMTC obtained in Chapter 4 are 30% higher in viscosity like water fluids in which oscillation flow is induced by segmentation.

On the other hand, when the viscosity of the solution increases by adding 1.0% (w/v) of guar gum, a similar back-and-forth pattern was observed. However, the velocity of the tracer due to segmentation was reduced by increasing the viscosity of the chyme. Figure 6.17 shows a characteristic flow pattern over time in the axial direction of the SIM. From Figure 6.17(b), it was possible to identify 10 sequential movements every minute. Peaks in the same ‘x’
direction are a result of the left contraction (cuff 1) while small peaks in the opposite direction of the x-axis, are produced by the right contraction of the SIM.

![Figure 6.17](image)

**Figure 6.17** Flow pattern of a radioactive tracer (600 µm) under segmentation contractions in the SIM for 1.0% (w/v) guar solution (a) 500-1000 seconds and (b) 60 seconds.

Assuming that the tracer is isokinetic with the fluid (density-matching must be tested for further experiments) the tracer patterns show that the strongest fluid motions are closed to the segmentation cuffs and two basic flow patterns are produced by the retropulsive back and
forth movements with the highest velocity up to 0.008 m/s. As previously found in PLIF experiments, contractions produced by the first cuff were more effective in terms of fluid motion than that produced by the second cuff. This could be explained by the restriction that the peristaltic pump imposed to the fluid movement.

Table 6.3 summarizes the mean flow velocities estimated from the experimental data obtained with PEPT.

<table>
<thead>
<tr>
<th>Guar gum (%, w/v)</th>
<th>Net flow</th>
<th>Local flow</th>
<th>Velocity (osc) m/s</th>
<th>Velocity (ave) (10^{-4}) m/s</th>
</tr>
</thead>
<tbody>
<tr>
<td>0.5</td>
<td>No net flow</td>
<td>2s</td>
<td>0.003</td>
<td>7.00</td>
</tr>
<tr>
<td>0.5</td>
<td>3 rpm</td>
<td>2s</td>
<td>0.050</td>
<td>9.10</td>
</tr>
<tr>
<td>0.5</td>
<td>3 rpm</td>
<td>1s</td>
<td>0.005</td>
<td>5.50</td>
</tr>
<tr>
<td>1.0</td>
<td>No net flow</td>
<td>2s</td>
<td></td>
<td></td>
</tr>
<tr>
<td>1.0</td>
<td>3 rpm</td>
<td>2s</td>
<td>0.007</td>
<td>2.70</td>
</tr>
<tr>
<td>1.0</td>
<td>3 rpm</td>
<td>1s</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Segmentation motion produces rhythmical contractions and relaxation of the intestine that mixing the lumen content. Flow profiles indicated that these concentric contractions produced micromixing of the digesta and as a consequence, mass transfer could be enhanced. In turbulent flow with Reynolds numbers > 2100, particles of digesta flow rapidly changing from one position to the other developed oscillatory movements or eddies showed in Figure 6.16. These eddies bring dissolved material to the intestinal wall increasing mass transfer processes. As eddies move rapidly, mass transfer in fluids with turbulent behavior is also rapid, much more so than that resulting from molecular diffusion in the unstirred laminar sublayer of the small intestine. Nevertheless, when viscous polymers are added to the food, mixing is impaired, retarding also absorption of actives.
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Although the SIM has proved that is able to reproduce the characteristic movements found in the small intestine, some considerations must be taken into account. There are limitations when blood pressure cuff in the SIM are used to reproduce segmentation. For example, the amplitude, pressure and frequency of contractions could vary compare to the real gut, resulting in different patterns of fluid motion dynamics.

On the other hand, the open-SIM was used through this chapter to be a more effective way of mixing because net and local flow are simulated in this configuration. On the contrary, the closed-SIM only allows local flow caused by segmental contractions. As both ends are closed, pendular movements cannot be simulated in the closed configuration, reducing mixing. Moreover, there is not net flow induced by the peristaltic pump due to both ends of the system are closed. This could impact on velocities and flow dynamics. Thus, regardless re-circulation, the open-SIM could be a closer representation of the real small intestine, in which longitudinal and radial mixing take placed due to retropulsion and recirculation flow patterns.

6.4 Conclusions

Flow patterns studied in this Chapter lead to the conclusion that segmentation produces little net flow of the fluid but it is a powerful medium of mixing in the small intestine bringing food particles to the radial axis of the lumen. Retropropulsive and oscillatory flows of the lumen content were generated during segmentation in the SIM. Flow vortices and eddies were identified as responsible for mass transfer enhancement.

Flow visualization techniques such as PLIF and PEPT shown to be useful techniques to describe flow fluid patterns in the small intestine as a result of mixing or segmentation.
contractions. Turbulent mixing was generated in solutions with low viscosity under segmentation. Nonetheless, mixing was impaired as the viscosity of the fluid increases.

The SIM is a realistic model that is able to predict digestion and absorption of foods in a physiological manner since it can reproduce in an accurate form the segmentation movements found in the small intestine. The SIM simulates the characteristic movement found in the small intestine with representative physiological features, volumes and flow rates. Although it has limitations, the SIM promises to be a powerful tool all in further studies of digestion and absorption.
CHAPTER 7 – ENCAPSULATION OF ACTIVES

7.1 Introduction

To impact on the delivery rate of active ingredients, structured and functional foods can be developed in a variety of ways. For example, as shown in previous chapters, by adding soluble fibres such as guar, pectin and CMC to model liquid foods, rates of digestion and absorption were significantly reduced. Although controversial, it is generally recognized today that dietary fibre plays a crucial role on postprandial glucose and insulin responses \textit{in vivo}. By changing the bulk viscosity of the intestinal fluid, the biopolymers will interact with each other because of entanglements creating a physical barrier for actives (e.g. glucose) crossing from the bulk to the intestinal wall.

Nonetheless, another area of opportunity to address dietary-related conditions such as obesity, hypertension and diabetes is building structures (encapsulated actives) capable to deliver molecules at the desired place and rate in the GI tract. Unlike viscous solutions in which no permanent bonds between individual biopolymers are formed, in encapsulated ingredients the biopolymers are cross-linked into a tridimensional structure. By encapsulating functional ingredients it could be possible, for example, to impact on satiety and control the release of active molecules such as glucose (Norton et al., 2006). Encapsulation also improves the stability of a bioactive compound by protection from factors such as pH, oxygen, UV light. As encapsulation can solve formulation problems, it is becoming very popular in the food sector for protection and sustainable release of actives.

This Chapter focuses on the behaviour of alginate gel particles and liposomes under simulated gastrointestinal conditions that can impact the rate of actives release \textit{in vitro} by
Chapter 7 – Encapsulation of actives

using a STR. As solid particles can also modify the flow properties of the chyme, the performance of the particles would also studied in the SIM as a function of mixing. Preliminary experiments were carried out for encapsulating glucose in alginate microspheres; however, the release of this sugar was fast due to the porosity of alginate beads and the molecular weight of glucose (MW 180). For that reason, riboflavin (MW 376) was selected as model to study the effect that encapsulation has on active release. Riboflavin and alkaline phosphatase (ALP) were encapsulated within alginate gels by ionotropic gelation (section 7.2) and liposomes (section 7.3) respectively, to produced micro- and nano- scale particles. Encapsulation efficiency and stability when stored at various pH and release of actives were some of the parameters evaluated to study whether these structures could be potential vehicles for the gastrointestinal delivery of bioactives. In addition, breakdown and release of actives was studied in the SIM as a function of 2s segmentation (see pattern in section 4.3).

7.2 Gelled particles

To form gel particles of riboflavin, aqueous solutions (5.0 ml) of alginate (1.0 - 3.0%, w/v) containing the vitamin (2.65 μM) were added dropwise into 20 ml solutions of calcium chloride (gelation medium) at various concentrations 0.1, 0.3 or 0.5 M. The solution was stirred gently under constant speed at room temperature. Agitation was left during 15 minutes to ensure gelation. Then, particles were filtered and washed with distilled water for further characterization. Table 7.1 summarizes experimental conditions used to encapsulated riboflavin. Since all the combinations were done, nine different formulations varying the concentration of alginate and calcium were tested.
Table 7.1 Experimental conditions used for building gel particles.

<table>
<thead>
<tr>
<th>Formulation</th>
<th>Alginate (% w/v)</th>
<th>CaCl₂ (M)</th>
<th>Cross-linking time (min)</th>
</tr>
</thead>
<tbody>
<tr>
<td>I</td>
<td>1.0</td>
<td>0.1</td>
<td>15</td>
</tr>
<tr>
<td>II</td>
<td>2.0</td>
<td>0.3</td>
<td>15</td>
</tr>
<tr>
<td>III</td>
<td>3.0</td>
<td>0.5</td>
<td>15</td>
</tr>
</tbody>
</table>

7.2.1 Gelled particles: size and entrapment efficiency (EE)

The resulting beads were spherical, with a mean diameter of 1.5 ± 0.3 mm. As table 7.2 shows the diameter of the beads decreased as the concentration of CaCl₂ increased, as a result of the cross links formed between calcium ions and alginate molecules. It is well known that during gel formation calcium alginate gels shrink resulting in loss of water (syneresis) and an increase in the concentration of the biopolymer. The structures produced with low concentrations of calcium (0.1 mM) were approximately 15% bigger than those elaborated with higher concentration of calcium ions reflecting a reduction of syneresis of the gel at low calcium concentrations. This is in agreement with results reported by Rayment et al. (2009).

The formulation of the microspheres produced and entrapment efficiency (EE) are also listed in Table 7.2. The EE of alginate gelled particles loaded with riboflavin was higher at low concentration of alginate and CaCl₂. For example, when 1% (w/v) of alginate was used the EE diminished from 88% to 77% when CaCl₂ concentration was increased from 0.1 to 0.5M. Overall as the concentration of alginate increased the EE decreased when compared at the same concentration of CaCl₂. These results are in agreement with those found by others authors (Rayment et al., 2009, Lee et al., 2008). The EE of gelled particles is usually reduced with increasing the concentration of calcium chloride in gelling medium and cross-linking time. This behavior could be explained as a denser and more compact structure was formed.
with less volume available for the entrapment of active substances. As a result of syneresis, harder and more organized networks were formed when gelled particles are exposed to high concentration of calcium ions during extended periods of time.

Table 7.2 Ca-alginate particles formulation and entrapment efficiency (EE)

<table>
<thead>
<tr>
<th>Assay</th>
<th>Alginate (% , w/v)</th>
<th>CaCl$_2$ (mM)</th>
<th>Diameter (mm)</th>
<th>EE (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>1</td>
<td>0.1</td>
<td>1.66 ± 0.03</td>
<td>88.09 ± 0.03</td>
</tr>
<tr>
<td>2</td>
<td>1</td>
<td>0.3</td>
<td>1.51 ± 0.09</td>
<td>86.56 ± 0.25</td>
</tr>
<tr>
<td>3</td>
<td>1</td>
<td>0.5</td>
<td>1.32 ± 0.02</td>
<td>77.45 ± 0.09</td>
</tr>
<tr>
<td>4</td>
<td>2</td>
<td>0.1</td>
<td>1.83 ± 0.04</td>
<td>86.96 ± 0.10</td>
</tr>
<tr>
<td>5</td>
<td>2</td>
<td>0.3</td>
<td>1.75 ± 0.10</td>
<td>86.78 ± 0.22</td>
</tr>
<tr>
<td>6</td>
<td>2</td>
<td>0.5</td>
<td>1.56 ± 0.06</td>
<td>86.60 ± 0.06</td>
</tr>
<tr>
<td>7</td>
<td>3</td>
<td>0.1</td>
<td>2.24 ± 0.19</td>
<td>85.36 ± 0.02</td>
</tr>
<tr>
<td>8</td>
<td>3</td>
<td>0.3</td>
<td>1.97 ± 0.08</td>
<td>84.98 ± 0.05</td>
</tr>
<tr>
<td>9</td>
<td>3</td>
<td>0.5</td>
<td>1.67 ± 0.13</td>
<td>83.83 ± 0.12</td>
</tr>
</tbody>
</table>

7.2.2 Particles morphology

The microspheres produced by ionotropic gelation were dried at room temperature and by means of a freeze-dryer, to evaluate the effect of drying process. Figure 7.1 shows scanning electron micrographs of the structure of gelled Ca-alginate particles dried at room temperature and by freeze drying. During air drying, the gel beads become smaller with dense matrices in which the active is embedded. Small fissures in the matrix surface were observed (Figure 7.1a). When the gelled beads were freeze dried, spherical porous matrices were formed which easily break upon handling. The size of freeze-dried particles remained almost the same as before drying while those of air-drying particles became smaller and denser. Drying method selection depends on the final application of these particles.
7.2.3 Release under simulated gastro-intestinal conditions

Release studies were carried out to examine the suitability of using calcium alginate matrix gel beads for actives control release. Release of actives was first studied in a STR simulating the physiological pH during gastric (pH=2.0) and intestinal (pH=7.0) digestion and then, using the SIM to have an approach of the release in vivo and the effect of mixing. The movements of the small intestine wall are considerably important on the absorption of the actives and their transport through the intestinal mucosa.

Microspheres from each formulation (1.0 g) were placed in 10 ml of simulated gastric fluid (SGF) (section 3.2.5) at pH 2.0. Vitamin (riboflavin) loaded microspheres were left in agitation for 2 hours at room temperature. After this time, the capsules were filtered and transferred into 10 ml of simulated intestinal fluid (SIF) pH 7 (section 3.2.5), trying to mimic the transition from the stomach to the intestine where they remained in agitation for 3 hours at room temperature. In order to determine the amount of vitamin released from the capsules in these two different environments, aliquots of 1.0 ml of the remaining solution under agitation were taken every hour during 5 hours for each experiment. The samples were then
analyzed with a UV/Vis spectrophotometer at 445 nm. Percentage of cumulative release was estimated to obtain the dynamic release profile.

The cumulative release expressed as percentage is plotted in Figures 7.2 and 7.3 according to alginate formulation. As figures show the rate of release was highly dependent of the concentration of both alginate and calcium concentration.

Figure 7.2a shows the release profile of the riboflavin entrapped within the 1.0% (w/v) alginate beads after being exposed to gastric pH (2.0) within 2 hours. The release of vitamin from the gel particles was very fast during the first 2 hours in gastric solution reaching up to 60% of the original concentration of vitamin. After this period under intestinal condition release gradually decreased reaching in most cases equilibrium.

The release of riboflavin from microspheres formed with 1.0% (w/v) of alginate into simulated gastric fluid at pH 2.0 shown up to 50% of release after 2 hours independently of the concentration of CaCl$_2$ in gelling media. However, as the pH increased microspheres produced with 0.5M CaCl$_2$ appeared to be more stables with minimum release. As expected, the cumulative release of vitamin increased as concentration of calcium ions decreased. This is thought to be caused by a reduction of Ca$^{2+}$-induced alginate cross-linking in the structure of the microsphere resulting maybe into bigger ‘pores’. Over time, disintegration of beads was observed with most of them breaking completely before the end of the experiment or in the worst scenario after exposure to simulated gastric fluid. Results from beads formed with 2% (w/v) of alginate are shown in Figure 7.2b. These trends were slightly different when compare with those for 1.0% (w/v) of alginate. In this particular case, microspheres were very unstable with almost 100% of the release of vitamin after 4 hours for formulations within 0.3 and 0.5M of CaCl$_2$. However, over time, disintegration of the beads was observed with one set of 1.0% (w/v) alginate beads breaking down completely before the end of the
experiment. This could be explained as a result of the increased concentration of monovalent ions in the solution.

![Graph showing the dynamic release of riboflavin from Ca-alginate beads into SGF and SIF at room temperature.](Image)

**Figure 7.2** Dynamic release of riboflavin from (a) 1.0 and (b) 2.0% (w/v) Ca-alginate beads into SGF and SIF at room temperature.
Dynamic release of vitamin from microspheres formed with 3% (w/v) of alginate is shown in Figure 7.3. When exposed to simulated gastric fluid, around 40% of the entrapped riboflavin was released during the first 2 hours for 0.1 and 0.5 M CaCl$_2$. Nevertheless for 0.3 M up to 75% had been released after gastric fluid exposure. Once placed in the intestinal solution, the rate of release was almost constant reaching equilibrium at 80, 75 and 50% for beads formed with 0.3, 0.1 and 0.5 M CaCl$_2$, respectively.

**Figure 7.3** Dynamic release of riboflavin from 3.0% (w/v) Ca-alginate beads into SGF and SIF at room temperature.

Overall, an inverse relationship between calcium ions concentration and vitamin release rates was found. As the concentration of calcium increases from 0.1 to 0.5 M the release of riboflavin was delayed. Thus, higher concentration of calcium ions in the gelling media induced a greater degree of cross-linking resulting in higher gel strength and limiting swelling patterns. This is agreement with results obtained by other authors (Rayment et al., 2009).
Regarding the concentration of alginate, all formulations follow similar trends; with a fast release during the first 3 hours, most notably when structures were placed into the simulated gastric solution at low pH. This suggests an initial weak structure of the bead possible due to dissociation of calcium ions at low pH (Rayment et al., 2009).

Alginate molecules are linear block co-polymers of β-D-mannuronic (M block) and α-L-guluronic acids (G block) with a variation in composition and sequential arrangements. When the alginate beads area placed in the gastric fluid (pH 2.0), the negatively changed carboxilate groups from uronic acids of alginate begin to protonate to form uncharged \(-\text{COOH}\) groups. This reduces the crosslinking due to decrease in ionic and electrostatic interactions. As a result, the bead structure becomes unstable with a subsequent faster release of encapsulated vitamin.

After 3 hours when the beads are transferred into the SIF at pH 7, the carboxylic groups of alginate are deprotonated \(-\text{COO}^\cdot\). Therefore, the beads begin to disintegrate along with the release of the encapsulated vitamin. In addition to this, the ion exchange between \(\text{Na}^+\) ions from the simulated fluids and \(\text{Ca}^{2+}\) ions from the egg-box cavities of the polyguluronate blocks of alginate also contributes towards the disintegration of alginate beads.

The ion exchange phenomena will impact not only on the swelling process and solubility of the particle but also the release of the active compound as a result of the surface area available for absorption. However, based on the results an increase in alginate concentration seems to improve the stability of the capsules under acidic conditions and i.e. towards swelling.
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From these figures, there is a clear effect on the interactions of alginate and calcium ions at different concentration, in terms of the encapsulation of the active and most importantly in the release of active components under simulated intestinal conditions. The encapsulation of vitamin B2 in Ca-alginate gels strongly depends on the cross-linking of calcium ions into the polymeric structure of the alginate. Thus, it would be recommended that the capsules were stable, dense and with high available volume to allow the entrapment of the active and therefore their resistance to the physiological conditions of the stomach (pH 2.0) to be released in the small intestine.

In conclusion, release tests performed simulating the GI pH variation also evidenced a direct relationship between alginate and CaCl2 concentration to the release of riboflavin; higher calcium concentration leads to a greater degree of cross-linking, thus should had resulted in a subsequent slowing of vitamin release rate.

Overall, the effect of cross-linking of the calcium ions with alginate plays an important role in the release of actives, the denser and more compact the capsule is, the smaller the pore of the beads are, therefore more resistance to the release at different values of pH. Although particles made with high concentration of alginate had a low EE, were more resistant to breakage in acidic conditions, so more appropriate for the release of actives in the intestine.

**7.3 Liposomes**

In this section liposomes were designed to protect the activity of active components from acidic condition in the stomach. As a model molecule in this particular case, alkaline phosphatase (ALP) was used and encapsulated within alginate gel core liposomes. Multilamellar vesicles (MLV) were prepared using the dry lipid film hydration by dissolving the enzyme in a solution of alginate and encapsulating this mixture in liposomes then
crosslinked the alginate by addition of CaCl$_2$ to the liposome suspension creating liposomes with a crosslinked gel core. These particles were characterised and compared to liposomes prepared without alginate (standard liposomes) using particle size analysis, ESEM and Cryo-SEM, entrapment efficiency, zeta potential and activity of the encapsulated enzyme following exposure to simulated gastric pH.

### 7.3.1 Liposomes: particle size and entrapment efficiency (EE)

Characterisation of the ALP loaded liposomes and gel core liposomes using laser diffraction revealed a large difference in mean particle size. The gel core liposomes were over twice the size of the standard MLV 30.8 ± 3.1 and 13.4 ± 0.1 µm respectively (Table 7.3). Entrapment however, showed no significant difference (p>0.05) between the vesicle types.

<table>
<thead>
<tr>
<th>Vesicle type</th>
<th>Vesicle size ± std (µm)</th>
<th>EE ± std (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>MLV Liposomes</td>
<td>13.4 ± 0.1</td>
<td>0.9 ± 0.3</td>
</tr>
<tr>
<td>Gel Core Liposomes</td>
<td>30.8 ± 3.1</td>
<td>1.6 ± 0.3</td>
</tr>
</tbody>
</table>

### 7.3.2 Liposomes morphology

As ESEM images shown, vesicle aggregation occurred in the gel core liposome sample (Figure 7.4a) with the individual particle size in the region of 10-15 µm which is in the same order of size as the standard liposomes, in which aggregation did not appear (Figure 7.4b). This is likely to be due to alginate present on the surface of the liposomes entangling with adjacent liposomes prior to crosslinking, then once crosslinking is initiated by addition of calcium, gelation occurs causing adhesion of the particles. The use of ESEM for studying changes in liposome morphology has been used previously which confirmed the utility of this
technique in providing a real-time alternative assay for liposomes stability and formulation (Mohamed et al., 2004).

![ESEM micrograph of (a) gel core liposomes and (b) conventional liposomes.](image)

**Figure 7.4** ESEM micrograph of (a) gel core liposomes and (b) conventional liposomes.

### 7.3.3 Liposomes: zeta-potential measurement

To determine whether alginate or ALP were present on the surface of the liposomes, measurements of zeta potential which is an indirect measurement of surface charge (Perrie and Gregoriadis, 2000) were performed on liposomes loaded with ALP and gel core liposomes loaded with ALP before and after crosslinking. These measurements were also performed on samples without ALP. Figure 7.5 shows that liposomes containing ALP and/or uncrosslinked alginate have a negative zeta potential however when the alginate is crosslinked the surface charge becomes positive. Interestingly the crosslinked liposomes containing ALP gave a significantly greater zeta potential ($p < 0.05$) than when there was no ALP encapsulated, $15.75 \pm 1.28$ and $12.38 \pm 1.38$, respectively.
These measurements support the presence of alginate on the surface of the vesicles as samples of gel loaded liposomes gave a negative zeta potential indicating the presence of negatively charged sodium alginate. When the gel loaded liposomes were crosslinked using calcium, the vesicles became positively charged due to the Ca$^{2+}$ ions suppressing the negative charge of the alginate. The presence of ALP which has an overall net negative charge (Beratis et al., 1970) affected the zeta potential on the standard liposomes which indicated that there may be some protein incorporated in the external lipid bilayer.

The binding of protein within the bilayers of liposomes is a well known phenomenon and occurs due to favorable electrostatic interactions between the protein and the phospholipid (Colletier et al., 2002). Surprisingly when ALP was incorporated with the crosslinked gel core liposomes there was an increase in zeta potential due to the overall net negative charge of the protein. ALP, however, is known to act as a calcium binding protein (de Bernard et al.,
1986) therefore addition of $\text{Ca}^{2+}$ as a crosslinker for the alginate, may have also been bound to the ALP shifting the overall charge of the ALP from negative to positive.

7.3.4 Liposomes internal structure: cryo-SEM analysis

Further evidence of alginate incorporated onto the surface of the gel loaded liposomes was revealed in the cryo-ESEM micrographs given in Figure 7.6a which shows the surface of a gel loaded liposome having small solid sections upon the surface thought to be crosslinked alginate. The presence of several thick layers is also evident when cross sectioned (Figure 7.6b) indicating gel is incorporated throughout the many lamellae of the liposome. Clusters of cross sectioned gel loaded liposomes are shown in figure 7.6c further indicating aggregation. Although it is clearly visible that the conventional liposomes are multilamellar as expected for MLV’s (Figure 7.6e) the aggregation characteristics and the thick layered lamella were not witnessed (Figure 7.6d-f) therefore it can be argued that incorporation of alginate alters the structural characteristics of liposomes potentially affecting the functional properties.

The presence of calcium alginate on the surface of alginate loaded liposomes has been hypothesised previously by Monshipouri and Rudolph (1995) who used differential scanning calorimetry to indicate a marked reduction in the enthalpy of the main gel to liquid crystalline phase transition of the lipid in MLV and LUV dispersions when alginate was encapsulated in the core (Monshipouri and Rudolph, 1995).
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Figure 7.6 Cryo-ESEM micrographs of (a) gel core liposomes before fracture, (b) cross-section of gel core liposome, (c) cluster of cross-sectioned gel core liposomes showing aggregation, (d) ALP-loaded liposome before fracture, (e) cross-sectioned ALP-loaded liposome with no evidence of aggregation.

7.3.5 Liposomes pH stability

The effects of pH on the stability of both vesicle types were analysed by monitoring vesicle size over a period of 10 days when incubated at pH 3.8 pH 7.4 and pH 10 (Figure 7.7). Standard MLV samples showed no dramatic change in mean diameter across the range of pH tested (Figure 7.7a). The gel core liposomes, however, appeared to decrease in size when stored at acidic pH and at pH 7.4 there was a substantial rise in mean diameter after 10 days storage (Figure 7.7b).

These results indicate that standard liposome samples remained stable over a range of pH (pH 3.8 - pH 10) perhaps showing slight instability at pH 3.8 highlighted by a slight increase in particle size (13µm to 15µm) (Figure 7.7a). The reduction in particle size of the gel core liposomes at pH 3.8 (below the iso-electric point of ALP but above the pKa the negatively charged carboxylate groups on the alginate) was possibly a result of the alginate gel contracting (Draget et al., 1994) as the pH approaches the pKa of the carboxylate group.
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When stored at pH 7.4 the increased size of the alginate containing liposomes following 10 days incubation may be due to the alginate gel gradually swelling. This 50% gain in size is similar to what was reported by Martins et al. (2007) for alginate beads.

![Figure 7.7](image)

**Figure 7.7** The effect of pH on the size of (a) MLV’s loaded with ALP and (b) gel core liposomes over a period of 10 days

### 7.3.6 Release of ALP

To determine the protective effect encapsulation in liposomes and gel core liposomes had on ALP, the loaded samples were exposed to simulated gastric pH (pH 2) for 2 hours then enzyme activity of the encapsulated ALP was evaluated in addition the quantity of ALP released by the liposomes and gel core liposomes was also monitored. The release curves in figure 7.8 reveal that the gel core liposomes released 13% of its entrapped ALP compared with 20% released from the standard liposomes over 2 hours. Multilamellar vesicles are expected to be more stable than other liposome types due to the many bilayers surrounding the encapsulated material (Kokkona et al., 2000) however the addition of alginate appeared to increase the stability in acid, where 7% less ALP was released from the gel core liposomes.
when incubated for 2 hours at 37 °C at pH 2. This stability was highlighted further by evaluating release over 48 hours at pH 2 which revealed the alginate containing liposomes retaining 70% ALP compared with 32% retained by the standard liposomes (results not shown), indicating the potential of these particles as sustained release delivery vehicles.

Figure 7.8 Release profiles of ALP from standard liposomes and gel core liposomes. Points represent mean values with vertical error bars indicating standard deviation, n=3.

7.3.7 Activity of ALP

Although it has been shown that ALP was successfully encapsulated and release in acidic pH was retarded by incorporation of alginate in the liposomes this does not confirm whether the ALP was still active. In addressing this, an activity assay was performed on the encapsulated APL during 2 hours exposure to pH 2. Figure 7.9 compares the activity of APL in the gel core liposomes with standard MLV and free ALP.

The activity of the free ALP was reduced on exposure to pH 2, however, both the MLV and the gel core liposomes protected the activity of the enzyme with no reduction in activity over the first 30 minutes of exposure. After 2 hours exposure activity was significantly higher in
the gel core liposomes (80% of original activity) than in the standard liposomes (55% of original activity) (p < 0.05).

![Figure 7.9](image_url)

**Figure 7.9** Percent of ALP activity retained during 2 hours of exposure to simulated gastric fluid, pH 2, for standard liposomes, gel core liposomes and free ALP.

The reduction in activity in the standard liposomes is a possible consequence of degradation of the liposomes or by acid ingress due to the pH gradient across the lipid bilayer. Moreover, evidence from the measurements of zeta potential (Figure 7.5) of ALP on the surface of the liposomes could also explain the reduction in activity. In the gel core liposomes, ALP activity is reduced over the 2 hours period but at a slower rate than was seen in the standard MLV. This can be explained by the acid insoluble properties of alginate gels. This acid insolubility however, is thought not to prevent a reduction in activity over longer periods of time as acid is gradually imbibed by the gel forming an alginic acid gel. This reduction in pH within the gel would ultimately reduce the activity of the enzyme.
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The tailorability of liposomes and biopolymers such as alginate provide scope for further investigation of these gel loaded particles as oral delivery vehicles as properties such as surface charge, particle size and stability are easily manipulated further by selecting alternative lipids to form the liposomes (e.g. cationic phospholipids), the addition of secondary processing steps such as extrusion and incorporating a bilayer stabilizer such as cholesterol.

7.4 In vitro release of gel particles and liposomes in the SIM

Release experiments were also carried out in the SIM to study the effect of mixing on the release of riboflavin and ALP, under physiological conditions. However, in the case of gel particles with a mean diameter of 1.5 mm, peristaltic pumps and the connectors used in the SIM, restricts its use to nanoscale particles. On the other hand, as concentration of riboflavin and ALP entrapped in gel particles and liposomes, respectively, was low, quantification of active released in the recipient side of the SIM (1.0 L) was very inaccurately. Liposomes for example, should be produced in large amounts to obtain representative results in the SIM. On the other hand, to measure ALP absorption in the recipient side of the SIM, the semi-permeable membrane must be changed because the molecular weight of the ALP is in the range of 140-160 kDa and the membrane has a MWCO of 8.0 kDa i.e. the enzyme cannot cross the permeable membrane used in this research.

7.5 Conclusions

To develop effective active delivery systems for controlled release, gel particles and liposomes were built for encapsulating riboflavin and ALP, respectively. Sodium alginate was used as a coating material. In both cases, alginate produced micro or nanocapsules that
protected the active ingredient from the acidic gastric environment. Gel particles were produced at various concentrations of alginate and CaCl$_2$.

The *in vitro* release study shows that microparticles of alginate have high pH sensitivity with a faster release under gastric conditions with a faster release at pH 2 (SGF). This behaviour was caused possibly by the surface area available for diffusion. The alginate structures were easily destabilized in the SGF. Overall, the release of encapsulated riboflavin decreased with increased capsule size and alginate concentration.

Alginate was also incorporated within the core of liposomes which dramatically affected liposomal characteristics both in terms of morphology and stability. Incorporating alginate into liposomes appears to improve stability at acidic pH however at neutral pH the alginate causes swelling and aggregation of the liposomes. This study also suggested that alginate was present on the surface as well as in the core of the liposomes. The alginate loaded liposomes offer potential as an delivery platform for functional proteins, verified by maintaining the 80% of the original activity of ALP following exposure to simulated gastric pH for a period of 2 hours.
CHAPTER 8 – CONCLUSIONS AND FUTURE WORK

This work focused on understanding mass transfer phenomena occurring during digestion and absorption processes in the small intestine from an engineering perspective. To predict the in vivo behavior of food digestion and actives absorption, a dynamic in vitro small intestine model was used. This versatile model reproduces anatomical features of the human small intestine simulating characteristic flow and mixing of the digesta by a unique pneumatic system. However, differences between the semipermeable membrane used in the SIM and the intestinal epithelial cell membrane must be taken into account when these processes are simulated.

On the other hand, it was found the configuration of the SIM (open or close) affects the rates of absorption. For glucose absorption and starch digestion, it was possible observed that the open-SIM with recirculation of the chyme, had a better performance in mimicking mixing in the small intestine than the close SIM. Since the latter restrict the movement of lumen content, longitudinal mixing was impaired, this was more evident when viscous solutions and starch food models were used. Another drawback of using the closed system is that the semi-permeable membrane could be easily damaged by the mechanical contractions applied to the SIM. Since the internal pressure of the lumen side of the SIM increased, the membrane could be submitted to mechanical deformation that can change its permeability i.e. driven to different results.

Basically, in this work two main points have shown; (i) how bulk-phase viscosity can have effects and (ii) how encapsulation of actives might work under gastrointestinal conditions. In both cases, a physical barrier was created to delay the rate of digestion and absorption of
active molecules. Some combination of the use of viscosity modifiers and encapsulated actives would be useful to control two parts of behavior.

8.1 Main conclusions

8.1.1 Actives absorption

Absorption of glucose was studied in the closed-SIM as a function of (i) mixing and (ii) food formulation. Results showed that mixing, induced by segmental contractions, increased the rate of glucose transferred by bringing material to the membrane of the SIM and i.e. to the epithelium of the small intestine \textit{in vivo}; reducing the unstirred layer formed near the membrane and helping diffusion of nutrients. Overall, mass transfer was enhanced when segmentation was applied to the system. It was found that when viscosity of the chyme increased, mixing and propulsion of the lumen contents were impaired, reducing the rates of glucose absorption.

On the other hand, food formulation has a significant incidence on glucose absorption and as a consequence on mass transfer. By increasing the viscosity of the digesta from 2 to 200 mPa.s, it was possible to reduce glucose absorption up to 90%.

Both mixing and food formulation have a significant effect on mass transfer phenomena in the SIM. Whilst mixing enhances mass transfer processes, increasing the viscosity of the food results in a reduction of the absorption across the membrane of the SIM.

In the second section, the open SIM configuration was used to study the effect that different biopolymers has on glucose absorption due to the viscosity of the solution and the frequency of segmental contractions (oscillation).
Chapter 8 - Conclusions and future work

The results showed that by adding soluble fibres such as guar gum, pectin and CMC on glucose solutions the rate of absorption i.e. mass transfer coefficients were slow down. On the other hand, wall movements influenced intestinal absorption. It was established that as the number (frequency) of segmental contractions decreased mass transfer coefficients were increased in viscous polymers such as CMC. However, frequency of the contractions does not seem to affect significantly the mass transfer coefficients in food models added with guar or pectin. This could be related to the shear thinning behaviour that each biopolymer has at higher shear rates.

CMC was more effective than other biopolymers and its performance was also affected by the frequency of contractions. This might relate to the shear thinning behaviour of CMC at high frequencies. According to the results the characteristic shear rate in the SIM can be set between 10 and 100 s\(^{-1}\). However, these values must be validated \textit{in vivo} by evaluating the performance of guar, pectin and CMC added to glucose solutions and monitoring the increase on blood glucose levels.

Thus, by understanding how polymers affect viscosity, novel functional foods and drugs can be designed to respond special needs of the consumer. On the basis of the results of this work it would be also possible to predict how other polymers might behave during intestinal digestion and absorption based on their chemical structure and rheological behaviour. These findings are very promising in designing diets for diabetic and obese people to cope with blood glucose levels.
Chapter 8 - Conclusions and future work

8.1.2 Starch digestion

For starch hydrolysis, the parameters investigated were degree of mixing by using a STR and the SIM, (ii) concentration of enzyme and substrate (starch); flow rate and food formulation in starch solutions and white bread.

It was observed that the degree of mixing impacted the performance and velocity of digestion. Compare to the STR a ‘lag time’ was observed in the SIM. The kinetics of starch hydrolysis in the SIM showed a sigmoidal behaviour with a characteristic ‘lag phase’ or ‘delay’ for both digestion of starch and absorption of glucose. Regarding to digestion, this behaviour is one of the main contributions of this work which is physiologically relevant and must be considered to further models development.

The rate of enzymatic reactions was reduced when guar was added delaying the access of actives to the absorptive epithelium due to a decrease of both propulsion and mixing. Similarly, this tendency was observed when bread digestion was mimicked in an aqueous solution (control) and in the presence of guar. When viscosity of the lumen content was increased, glucose absorption was reduced up to 45%. Compare with glucose and guar, it was observed that absorption was faster in a model glucose system in which no chemical reactions take place before absorption. On the other hand, absorption was delayed due to the presence of starch that increases the viscosity of the bulk lumen.

When white bread was tested in the presence of guar, absorption decreases. Consequently, viscous solutions impaired both enzyme-substrate binding and glucose diffusion from the bulk to the wall of the lumen side of the SIM.
By using the SIM it was demonstrated that increasing the viscosity of food digesta resulted in a reduction of concentration of nutrients available for absorption. Increasing the viscosity of the lumen or biopolymer side resulted in a decrease of the mass transfer coefficients.

The model provides an accurate and meaningful method for predicting the performance \textit{in vivo} of compounds and nutrients in the small intestine. The food formulation, starch and enzyme concentrations are key factors to control digestion of foods during in vitro digestion that should be taken into account in designing in vitro GI tract models.

\subsection*{8.1.3 Mixing processes}

Since mixing processes are crucial for chemical reactions and absorption processes in the GI tract. Flow visualization techniques were employed to understand the impact that segmentation has on mass transfer in the gut. It was hypothesized that mass transfer in the lumen side was enhanced by formation of eddies or recirculating flow as a result of mixing. Furthermore, mixing brings material to the intestinal wall reaching the absorptive sites present in the gut.

Based on the results, it was concluded that segmentation produces little net flow of the lumen content but promotes radial mixing. It was found that contractions applied in the SIM are a powerful medium of mixing in the small intestine, bringing food particles to the wall of the lumen. Retropropulsive and oscillatory flows of the lumen content were generated during segmentation. Flow vortices or eddies were identified as responsible for mass transfer enhancement in the SIM. Turbulent mixing was generated in solutions with low viscosity under segmentation. Nonetheless, mixing was impaired as the viscosity of the fluid increases.
Flow visualization techniques such as PLIF and PEPT shown to be useful techniques to describe flow fluid patterns in the small intestine as a result of mixing or segmentation.

8.1.4 Encapsulation

Novel structures were developed for sustained release of actives. Riboflavin was encapsulated in gel microspheres while liposomes were used to encapsulate ALP.

The stability of alginate beads loaded with riboflavin were affected by both concentration of CaCl₂ and sodium alginate. The encapsulation efficiency was higher as the concentration of calcium decreased creating structures with high water content. Even though the efficiency of encapsulation was low for beads with high concentration of CaCl₂, these microspheres were more resistant to the simulated acidic conditions of the stomach pH 2. Those can be used to delivery of actives in the intestinal conditions due to its stability.

Incorporating alginate into liposomes appears to improve stability at acidic pH, however, at neutral pH the alginate causes swelling and aggregation of the liposomes. This study also suggested that alginate was present on the surface as well as in the core of the liposomes. The alginate loaded liposomes offer potential as an delivery platform for functional proteins, verified by maintaining the 80% of the original activity of ALP following exposure to simulated gastric pH for a period of 2 h.

By understanding how food is processed in the GI tract it would be able to design and develop structured food to manipulate digestion, reducing and delaying absorption of actives. Understanding how food breaks down during digestion and absorption processes can be applied to designing foods specifically targeted to a particular consistency and composition of
the bolus. The results showed that adding soluble fibres such as guar gum, pectin and CMC in glucose solutions affected rate of absorption i.e. mass transfer coefficients delaying the concentration of glucose. Then, foods can be structured in a variety of ways and each structure can have a different physiological response.

The SIM is a realistic model able to predict digestion and absorption of foods in a physiological manner since it reproduce in an accurate form the segmentation movements found in the human small intestine. This *in vitro* model simulates the characteristic movement found in the small intestine with representative physiological features, volumes and flow rates. Although it has limitations, the SIM promised to be a powerful tool in further studies of digestion and absorption.

### 8.2 Future work

The future work could be potentially done in various areas:

**Model validation.** A proper validation must be done in order to correlate the results obtained in the SIM with *in vivo* studies. Experiments by using MRI or visualization techniques could help to monitor contractions (amplitude, frequency) occurring in the small intestine and to evaluate the flow and concentration of enzymes as chyme enters to the gut. To avoid complications of further metabolism/clearance of glucose from the circulation, it is proposed to measure glucose in the hepatic portal vain rather than the periphery.

**Different biopolymers** such as β-glucans and fructans e.g. inulins at various concentrations and frequencies of segmentation could be also tested in the SIM. It has been stated recently that those biopolymers have a great impact on reducing glucose and cholesterol levels (Anttila et al., 2004). The SIM has been proved to be a useful tool to predict the performance of a biopolymer in the gut it could be interesting to evaluate the effect of other soluble fibres.
Chapter 8 - Conclusions and future work

**Digestion of complex foods**, including additional points of segmentation. This work was focused on model viscous fluids from either starch or glucose. However, real foods are very complex structures in which macromolecular interactions must be taken into account when processes of digestion and absorption are being modeled. There is also crucial to add not only amylases but also proteases and lipases for proteins and lipids digestion, respectively.

Study of **encapsulated particles** in combination with the bulk viscosity of the SIM. Although the design of the SIM allows to test different semisolid foods, the study of encapsulate particles in this model has limitations for the use of peristaltic pumps that can break those particles before they arrive to the intestinal side of the SIM. As such the mechanical system of the SIM can be modified to test the release of micro or nanostructures on simulated intestinal conditions. Structures breakdown can be also evaluated in the SIM.

**8.2.1 Absorption studies**

The present work was limited at three different biopolymers (guar, pectin and CMC) and frequencies (1s, 2s and 3s), however, further experiments can be done by using additional soluble and insoluble fibres and by changing the sequence of segmental contractions in a physiological context. For example, the maximum delay time between contractions is fixed at 6s (Labview software) but it could be possible to extend until ten seconds modelling a slow transit and less effective degree of mixing that could be physiological relevant in some real cases. The SIM could be also used along with other models such as Caco-2 to evaluate permeability of actives to the cellular membrane. By combining both models, complementary results can be obtained i.e. limitations of the SIM could be overcome. Caco-2 cells are able to mimic active transport absorption process.
Moreover, additional segmentation points can be included to the model by implementation of three or more inflatable cuffs. As such the length of the membrane has to be increased and maybe food recirculation could be avoided. To simulate motility in the human small intestine a period of no contractions, long period of contractions and regular contractions can be also simulated. One of the advantages of the SIM is that a variety of conditions can be simulated.

8.2.2 Digestion studies

Related to digestion studies, various starch types could be evaluated in the SIM. For instance, starches with different crystallinity or resistance to amylolytic attack (resistant starch) can be testing. Interactions among macronutrients (proteins, carbohydrates and lipids) must be also assessed. In this work, homogeneous solutions of starch were used, nevertheless foods have heterogeneous structures, thus for further experiments it is suggested to study complex foods and the interactions between ingredients. Multiphase meals, e.g. emulsions, can be also used to evaluate interactions among food components. Moreover, a mixture of proteases and lipases must be employed in combination with amylases to represent physiological relevant conditions in the small intestine.

Improvements in the method to feed the enzyme in the lumen side of the SIM must be done, including additional sampling points inside of the membrane during experimentation, taking into account that the concentration of the lumen changes with time and position, driven to a better understanding of the kinetics in the small intestine.

It is also possible to include additional steps to simulate digestion in the mouth and the stomach
Chapter 8 - Conclusions and future work

8.2.3 Flow visualization studies

Improvements in this area could be done by analysing quantitatively velocities generated in the SIM as a function of mixing and using another optical imaging techniques such as Particle Image Velocimetry (PIV). These studies must be done in combination with mathematical modelling experiments to predict velocity patterns and their influence in digestion and absorption processes.

A quantitative analysis must be done to calculate velocities and the trajectory of the particle base on the position of the tracer from PEPT results.

8.2.4 Encapsulation studies

Different encapsulation techniques and materials can be used in the future to evaluate the effect of microparticle structure in the release of actives and absorption. Structures, for example, unilamellar liposomes could be made to increase the entrapment efficiency of the internal structure. Moreover, starch could be encapsulated to delay its digestion and further glucose absorption. Due to time restrictions, further evaluation of the release and diffusion of encapsulated actives in the SIM cannot be done in this work. Analysis of the performance of encapsulated active release in the in vitro model to measure diffusion and mass transfer coefficients is recommended. In addition, combination of bulk viscosity and encapsulated actives could be done to evaluate the effect of both.

Validation of the SIM must be an integral part in future works to improve or change not only mechanical features of the system but also physiological concentrations of enzymes pH, T, residence time and secretions in the lumen side of the SIM.
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APPENDIX

Published work from this thesis (Proceedings)


Appendix I

ABSTRACT

The main objective of this work was to understand phenomena occurring during food digestion by using a dynamic in vitro Small Intestine Model (SIM), of particular interest was to study the effect of mixing and food formulation on starch hydrolysis and glucose absorption. The SIM reproduces the characteristic mixing of the human small intestine by expansion and contraction of two cuffs around the external tube. Corn starch hydrolysis was studied in single phase fluids varying the concentration of amylase and the viscosity of digesta by adding guar gum (0.5%, w/v) under simulated physiological conditions.

Results showed the effect of segmentation motion on nutrient delivery to the intestinal wall as a consequence of changes in the mass transfer coefficient. This is most likely due to the increased mixing in the SIM. The rate of glucose absorption was decreased from 0.75 to 0.48 µM/s when guar gum (0.5%, w/v) was added to the corn starch solution. As the viscosity of the solution increased from 2 to 200 mPa.s glucose absorption was reduced up to 50%. Experiments of starch digestion in the SIM, with and without the presence of guar, have shown that guar reduces the rate of starch digestion by impairing mixing and reducing diffusion within the fluid.

This research has demonstrated the capability of using an in vitro rig to simulate and obtain an engineering understanding of transport phenomena occurring in the small intestine leading to absorption of active components. Current work is aimed at developing a reaction engineering understanding of the gastrointestinal tract.

Keywords: Starch hydrolysis; digestion; absorption; glucose; small intestine.

INTRODUCTION

When refined sugars and rapidly digested carbohydrate are included on a regular basis in our diets, blood glucose and insulin responses are usually enhanced, resulting in dietary related chronic diseases such as diabetes and obesity [1-4]. To avoid this scenario, structured foods with specific health benefits should be designed in conjunction with studies to understand how food formulation impacts on digestion and absorption rates in the gastrointestinal tract (GIT).

Digestion is a very complex process at which the small intestine is a key step responsible for most of the active molecules absorption. To understand and reproduce this process, in vitro and in silica models have been designed [5-7]. Nevertheless, details of flow dynamics and mixing have not been extensively studied. Mixing of lumen contents is essential in enhancing food digestion and absorption. Intestinal motility allows interactions between food and pancreatic digestive enzymes and biliary salts that enable food digestion. Intestinal absorption is also influenced by the physical form of the food ingested and the physical properties (viscosity, density, concentration, flow rate) of the lumen content. It has been reported that increasing viscosity of the lumen content, for example, could reduce the nutrient absorption through the epithelium reducing glucose levels in blood [2,3]. Addition of viscous fibers into the food not only increases the viscosity of the lumen but also may protect starch from enzymatic attack. Overall, by changing food formulation, the rate of nutrients absorption could be modified resulting in structured food with a health benefits for the consumer.

The main objective of this work was to reproduce and model starch digestion by studying the effect of mixing and viscosity on glucose absorption rate. Digestion and absorption were studied by using a dynamic small intestine model simulated the physiological conditions of the human small intestine.
MATERIALS & METHODS

All reagents and digestive enzymes were purchased from Sigma-Aldrich (UK). Pancreatic α-amylase from porcine (30 units/mg solid SIGMA-Aldrich, UK) and amyloglucosidase from Aspergillus niger (300 units/mL SIGMA-Aldrich, UK) were used for starch digestion. Cornstarch (1.0%, w/v) was previously gelatinized in a boiling water bath during 30 minutes with intermitted mixing. After gelatinization, the starch solution was cooled at room temperature. The enzyme solution (50 ml) was freshly prepared for the digestion analysis by dissolving the pancreatic α-amylase (2.5 and 25 U/ml) and amyloglucosidase (3 U/ml) into a pancreatic mix solution for 10 minutes at room temperature. The Small Intestine Model (SIM) used in this work was designed to represent the characteristic flow and mixing processes in the small intestine by expansion and contraction of two cuffs around the external tube (Fig. 1b). This model functions as a concentrically mass transfer exchanger composed of an inner semi permeable membrane of cellulose (Spectra/Por 7®, MWCO 8000, 3.2 cm of diameter, Medicell International Ltd London, UK) and an outer non-active and impermeable tubing (5 cm of diameter, Flexihose, UK). The reaction was started by pumping the pancreatic solution into the SIM at 0.3 and 3.0 ml/min (Figure 1).

Glucose concentration was monitored in the lumen (inner tube) and recipient side (outer tube) for the control and when the viscosity of the digesta was increased by adding guar gum 0.5% (w/v). The concentration of glucose was calculated following the method of 3,5-dinitrosalicylic acid (DNS) reagent for reducing sugars. All experiments were performed at least in triplicate. The rate of starch digestion in the lumen side and glucose absorption in the recipient side, were obtained from the slope (mM/s) of each plot by a linear regression analysis. Overall mass transfer coefficients (K) were estimated by Eq. 1.

\[ N_i = K(c_i - c_\infty) \]

where \( N_i \) is the molar flux (mol/m²s) including both diffusion and convection and \( c_i \) (mol/m³) is the concentration at the interface of the same fluid as the bulk concentration \( c_\infty \) (mol/m³).

RESULTS & DISCUSSION

Food digestion was done by using an in vitro dynamic small intestine model that simulates representative flow and mixing processes in the small intestine. The effect of mixing and food structure was studied on the glucose absorption phenomena. Figure 2 shows the data from the starch digestion in the lumen side (inner side or membrane) and the glucose absorption after generation in the recipient side (outer side) in the SIM varying the concentration of amylase per minute. An increase of glucose concentration was observed in both sides of the SIM as the concentration of amylase was increased over time. Starch hydrolysis was rapid rate in the first minutes of the enzymatic reaction following by a progressive decreased rate, reaching the steady state after approximately 60 min according to the experimental conditions with maximum glucose concentration up to 40 mM. Glucose absorption was lineal in the recipient side with a maximum of approximately 5 mM, almost ten-fold less than the glucose released from the lumen as a result of the enzymatic starch digestion.
Figure 2. Digestion in the SIM, (a) starch digestion in lumen and (b) glucose absorption in recipient side.

Table 1 shows the rate of digestion of starch and glucose absorption calculated from a linear regression of the data plot in Fig. 2. The Overall Mass Transfer coefficients were estimated from the flux of the lumen to the recipient side with the concentration difference at the maximum concentration reached at the steady state in the lumen.

Table 1. Digestion and absorption rates for the experimental conditions.

<table>
<thead>
<tr>
<th>Experiment</th>
<th>Digestion (µM/s)</th>
<th>Absorption (µM/s)</th>
<th>Overall Mass Transfer Coefficient (µm/s)</th>
</tr>
</thead>
<tbody>
<tr>
<td>2.5 U at 0.3 ml/min</td>
<td>6.91 ± 1.23</td>
<td>0.40 ± 0.04</td>
<td>0.33 ± 0.03</td>
</tr>
<tr>
<td>25.0 U at 0.3 ml/min</td>
<td>9.22 ± 2.34</td>
<td>0.49 ± 0.21</td>
<td>0.35 ± 0.09</td>
</tr>
<tr>
<td>2.5 U at 3.0 ml/min</td>
<td>8.43 ± 0.78</td>
<td>0.51 ± 0.01</td>
<td>0.38 ± 0.05</td>
</tr>
<tr>
<td>25.0 U at 3.0 ml/min</td>
<td>54.55 ± 2.11</td>
<td>0.48 ± 0.05</td>
<td>0.30 ± 0.02</td>
</tr>
</tbody>
</table>

The rate of starch hydrolysis increased up to 90% as the concentration of enzyme increased ten-fold in the lumen side, reaching the maximum value at 25.0 U and 3.0 ml/min while the minimum was observed for the lowest concentration of enzyme at 2.5 U and 0.3 ml/min. Results in the lumen also indicated that there was no significant difference between the starch hydrolysis rates when the concentration of enzyme per ml of digesta was identical. This implies that no matter how fast or concentrated the pancreatic solution is feeding into the SIM if the units of enzyme per time per volume are the same, the hydrolysis will take place at similar rates of reaction in well-mixed solutions with viscosities like water.

The increase of enzyme concentration per ml of digesta markedly increased the hydrolysis of starch but had little effect on glucose absorption (Table 1). The glucose profiles in the recipient side appeared to be linear. No appreciable amounts of glucose were detected at the beginning of the experiment. From the highest enzyme concentration, the change in glucose was detectable at 30 min into the recipient. However, there was a “delay” in glucose absorption (from 30 to 60 min) when the rate of starch digestion decreased. Once glucose was generated and released from the starch structure in the lumen side, absorption took place. Interestingly, this could be correlated to the point where viscosity of the starch solution falls in the lumen.

Effect of food formulation

In Figure 3 glucose absorption data are shown for products having different viscosities in the lumen side. Absorption appeared to be linear (R² of 0.95) resulting in a gradient of 7.45 x 10⁻³ mM/s. The flux, N, was calculated to be 1.55 x 10⁻⁵ mol/m²/s. By dividing the molar flux by the concentration difference in mmol/m³ an overall mass transfer coefficient, K, was obtained as 4.83 x 10⁻⁷ m/s for the aqueous starch solution and 3.10 x 10⁻⁷ m/s for the starch solution added with guar both under segmentation conditions.
Figure 3. Glucose absorption levels in the side of the SIM, with (0.5%, w/v) and without guar gum for two food models (a) corn starch and (b) white bread.

The effect of the presence of guar was shown by 36% reduction in the mass transfer coefficient when mixing was applied (Table 2). The data not only suggest that the rate of enzymatic reactions was reduced when guar was added but also that viscous solutions can delay the access of actives to the absorptive epithelium due to decrease of both propulsion and mixing [2]. Similarly, this tendency was observed when bread digestion was mimicked in an aqueous solution (control) and in the presence of guar. When viscosity of the lumen content was increased from 2 to 200 mPa.s, glucose absorption was reduced almost 15% after one hour, however, no significant difference was observed at the first minutes of digestion (Fig 1b). While the overall mass transfer coefficient from bread with guar gum was reduce 20%. These results are in agreement to those reported for Tharakan et al. [7].

Table 2. Rates of absorption and mass transfer coefficients for two food models.

<table>
<thead>
<tr>
<th>Experiment</th>
<th>Absorption (µM/s)</th>
<th>Overall Mass Transfer Coefficient (µm/s)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1. Starch solution</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Control</td>
<td>0.75 ± 0.05</td>
<td>0.48 ± 0.05</td>
</tr>
<tr>
<td>Guar gum (0.5%, w/v)</td>
<td>0.48 ± 0.02</td>
<td>0.31 ± 0.08</td>
</tr>
<tr>
<td>2. Bread</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Control</td>
<td>0.67 ± 0.01</td>
<td>0.43 ± 0.04</td>
</tr>
<tr>
<td>Guar gum (0.5%, w/v)</td>
<td>0.50 ± 0.02</td>
<td>0.32 ± 0.02</td>
</tr>
</tbody>
</table>

Overall mass transfer coefficient was reduced when guar gum (0.5%, w/v) was added to the solution of starch and bread. That could be explained by inhibition of propulsion and mixing in the membrane making interactions between the starch and amylase also difficult. Thus, starch digestion was lower in the presence of guar. Results suggested that molecular delivery is largely influenced by the fluid dynamics of the lumen side. Viscous polymers create a physical resistance for actives release and delivery that could modified the rate of nutrients absorption for the design of healthy diets with specific targets.

From the starch digestion experiment it was possible to show the effect that segmentation motion and food formulation had on nutrient delivery to the intestinal wall as a consequence of changes in the mass transfer coefficient. On the other hand, the rate of food digestion might be important in assessing the extension at which they raise the blood glucose levels in normal and diabetic people [8].
ACKNOWLEDGMENTS
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CONCLUSION
This research has demonstrated the capability of using an in vitro rig to simulate and obtain an engineering understanding of transport phenomena occurring in the small intestine leading to absorption of active components. Current work is aimed at developing a reaction engineering understanding of the GIT. Using an in-vitro system it was demonstrated that increasing the viscosity of food digesta results in a reduced the concentration of nutrients available for absorption.

REFERENCES
Appendix II

An in-vitro small intestine model for studying absorption of encapsulated actives

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A fundamental understanding of the performance of food structure in vivo will enable to design formulations that will provide specific health benefits to the consumer. In order to achieve this, an in vitro small intestine model (SIM) was designed and implemented to reproduce segmentation, which provide most of the mixing. The SIM was used to study the effect of formulation in the absorption rate. A range of biopolymers was used: CMC and guar gum (0 - 0.5%, w/v). The effect of formulation on actives from particulate delivery systems was also investigated, using gelled particles. Results showed that absorption of actives depends on the viscosity of the fluid. Fluids containing biopolymers exhibited slower absorption rates. Similarly, use of particulate systems demonstrated a significant effect in the delivery rates. The entrapment efficiency of calcium alginate beads was reduced with increasing the concentration of calcium ions in gelling medium and cross-linking time. Overall this work demonstrates the capability of using an in-vitro rig to simulate and obtain an engineering understanding of transport phenomena occurring in the small intestine leading to absorption of active ingredients.

Keywords: absorption, small intestine, encapsulation, structured foods

1 INTRODUCTION

Small intestine is a crucial step in digestion and absorption of essentially all dietary organic molecules. Despite of its importance in the delivery of active molecules relatively little work has been done to understand and quantify mass transfer phenomena inside the gastrointestinal tract (GI tract). In an effort to reproduce the diffusion and absorption processes in the gut, in-vitro and computational models have been developed [1-3]. However, just a few of them highlighted the importance of mixing in the small intestine. The mixing or segmentation contractions occur when a section of the small intestine becomes filled with digesta; as a result a localized concentric contraction takes place in the gut [4]. This process is repeated at spaced intervals along the intestine and is responsible for the fluid dynamics in the small intestine.

Intestinal absorption is not only related to the flow and mixing of fluids in the gut but also to the structure of the food and drug systems. For instance, it has been demonstrated that adding guar gum, pectin and fibers in a food system can notably delay absorption and transit time of glucose [5,6]. The rate of intestinal digestion, absorption and transit of the food could be also influenced by the physical form of food ingested and the physical properties of the intestinal contents produced [7,8].

Overall, the food industry needs to develop foods that deliver active ingredients with specific functions over an extended period of time. Potentially, this could be accomplished developing encapsulated structures which can be released in the right place and time.

The objective of this research was to understand absorption process and the effect of formulation in an in-vitro small intestine model.

1.1 Description of the small intestine model

Small intestine is comprised by the duodenum, jejunum, and ileum, reaching a length between 2 and 6 m [4]. The external diameter of the gut is on average 3.5 cm, varying from 5 cm in the upper portions of the duodenum to less than 2 cm in the distal ileum [1].

The in-vitro small intestine model (SIM) used in this research was designed to represent as closely as possible the human small intestine with a mean diameter of 3.2 cm and a characteristic length of 50 cm (instead of 6 m). The SIM functions as a concentrically mass transfer exchanger composed of an inner semi permeable membrane of regenerated cellulose (Spectra/Por 78, MWCO 8000, 32 mm of diameter, Medicell International Ltd London, UK) and an outer non-active and impermeable membrane (5 cm of diameter). This in-vitro model, gives an accurate representation of the flow and mixing processes in the small intestine [7].

These processes are simulated by the inflation and deflation of two cuffs (C1,2) around the external tube by alternatively applying compressed air (CA) and a vacuum (P2) (Fig. 1). The mechanical squeezing mechanism and the delay times are controlled by a computer, allowing the opening or closing of four different solenoid valves (V1,4) (Fig. 1).

The characteristics of the intestinal cell set up were: length of the mass transfer cell 50 cm (dialysis membrane), cuff length 12 cm, diameter of the inner tube 3.2 cm and diameter of the outer tube 5 cm.
2 MATERIALS AND METHODS

Glucose, Guan gum (GG), Carboxymethyl cellulose (CMC), methylene blue and sodium alginate were supplied by Sigma-Aldrich Company Ltd. (UK).

2.1 Absorption of actives in the SIM

A series of experiments were carried out using the SIM and glucose as a model system. Two different biopolymers, GG and CMC, were assayed. The biopolymer and glucose (55 mM) were homogeneously dispersed in distilled water using a magnetic stirrer. Two processing conditions were investigated according to the method used by Tharakan et al. [7]: (i) Net flow induced by a peristaltic pump (P1 in Fig. 1, 120 rpm) and no segmentation movements (NS), (ii) Net flow induced by peristaltic pump (P1 in Fig. 1, 120 rpm) and local flow from segmentation (SM) by cuffs C1 and C2 (Fig. 1) alternatively with a 2 second inflation time, 2 second deflation time, 2 second delay and 0.55 barg inlet air pressure. Both experiments were carried out in triplicate. Diffusion across the membrane was monitored at time intervals during one hour by measuring glucose concentration in the recipient according to the method of DNS for reducing sugars [9].

2.2 Mass transfer coefficients

Overall mass transfer coefficients (K) were estimated by Eq. 1.

\[
N_1 = KA(c_i - c_w)
\]

Where \(N_1\) is the molar flux (mmol/m² s) including both diffusion and convection and \(c_i\) (mol/m³) is the concentration at the interface of the same fluid as the bulk concentration \(c\) (mmol/m³). The reciprocal of the overall mass transfer coefficient \(K\) can be represented by the sum of the film resistances and that through the membrane and the water side (Eq. 2).

\[
\frac{1}{K} = \frac{1}{k_{bp}} + \frac{\frac{l_{mem}}{D_{mem}}} + \frac{1}{k_{rep}}
\]

Where \(k_{bp}\) is the biopolymer side mass transfer coefficient, \(D_{mem}\) is the diffusion coefficient of nutrient through the membrane, \(l_{mem}\) is the thickness of the membrane and \(k_{rep}\) is the recipient side mass transfer coefficient.

To find the mass transfer enhancement from mixing it is important to calculate the Sherwood number (Eq. 3).

\[
Sh = \frac{k_{bp}L}{D}
\]

Where \(k_{bp}\) is the biopolymer side mass transfer coefficient, \(L\) is the length scale (0.032 m) and \(D\) is the diffusion coefficient of glucose (6.9 x 10⁻² m²/s).

2.3 Encapsulation of actives

Sodium alginate (3.0%, w/v), glucose (10%, w/v) and methylene blue (0.005%, w/v) as an indicator were homogeneously dispersed in distilled water by means of a magnetic stirrer. This solution was dropped using a syringe (0.7 mm of inner diameter) into a second solution, containing CaCl₂ (1.0 and 5.0%, w/v) under gentle stirring. Microspheres (MS) were instantaneously produced by ionotropic gelation. The MS formed were allowed to stand in the solution for 5 and 10 minutes and then separated from the reaction mixture by filtration and washed with distilled water.

2.4 Entrapment efficiency

An indirect method was used to quantify the amount of colorant in the MS [10]. Aliquots from the filtered solutions remaining after removal of the beads were assayed using a spectrophotometer (CECIL instrument CE 1020) at 665 nm. The amount of substance entrapped was calculated from the difference between the total amount of the colorant added and the amount found in the filtered solution.

2.5 Release of actives

For this study calcium alginate beads were made according to the method previously described. The assay was done using a CaCl₂ solution (1.0%, w/v) and the cross-linking time was 5 minutes. After filtration the alginate beads (3.5%, w/v) were placed in 1000 ml of distilled water and added into the SIM. The experiment was done applying segmentation motion to study the effect of the mixing in the release of glucose and methylene blue. The amount of colorant released was calculated using a spectrophotometer at 665 nm (maximum absorption).

3 RESULTS AND DISCUSSION

3.1 Absorption of actives

In Fig. 2 typical data is shown for concentration versus time for guar gum and CMC when segmentation movements (SM) and no segmentation (NS) were applied. According to the results showed in Fig. 2, glucose mass transport process was accelerated when segmentation
movements were applied. After one hour, glucose concentration was around 6 mM when segmentation was induced in the SIM and 5 mM without segmentation for GG. This behavior was more evident for CMC at higher viscosities than GG (Fig 2). It possibly due to the increased mixing and a reduced boundary layer at the membrane surface [7].

Figure 2. Glucose increase in the SIM using Guar Gum (0.1%, w/v) and CMC (0.1%, w/v).

The overall mass transfer coefficient K (Tab. 1) was estimated for the range of biopolymers used. These results show the effect of changing the formulation of the system by increasing the viscosity. As the viscosity of the biopolymer was increased the absorption of glucose was reduced and the effect of segmentation motion was increased.

Table 1. Overall mass transfer coefficients in the SIM from different viscous materials.

<table>
<thead>
<tr>
<th>Biopolymer</th>
<th>Viscosity (Pa.s)</th>
<th>K (10^-7 m/s)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Control</td>
<td>0.001</td>
<td>5.35 ± 0.07</td>
</tr>
<tr>
<td>GG_0.1%</td>
<td>0.002</td>
<td>5.31 ± 0.85</td>
</tr>
<tr>
<td>CMC_0.1%</td>
<td>0.025</td>
<td>4.55 ± 0.05</td>
</tr>
<tr>
<td>CMC_0.5%</td>
<td>0.200</td>
<td>3.41 ± 0.37</td>
</tr>
</tbody>
</table>

The maximum overall mass transfer coefficient was found for water (control) under segmentation condition. This was expected as in this low viscosity solution the resistance to mass transfer from the bulk of the fluid to the surface of the membrane is minimum. The overall mass transfer coefficient decreased approximately 15% when viscosity of the fluid increased ten-fold. However, when viscosity increased from 0.001 Pa.s (water) to 0.2 Pa.s (CMC, 0.5%) K was reduced up to 90%. Overall resistance of the system to mass transfer was found to be 1.86 x10^6 s/m. For the fluids containing biopolymer k_{op} was estimated using (Eq. 2). The range of mass transfer coefficient in the biopolymer and relevant Reynolds numbers are shown in Tab. 2 when segmentation movements were applied. The change is due to the variation in the apparent viscosity from each biopolymer used (Tab. 1). The maximum velocity (0.06 m/s) was estimated using the displaced volume moved by segmentation motion. The diameter of the membrane was 0.032 m.

Table 2. Mass transfer coefficients and Reynolds numbers of the biopolymer side from different viscous materials.

<table>
<thead>
<tr>
<th>Biopolymer</th>
<th>K_{op} (10^{-7} m/s)</th>
<th>Re</th>
</tr>
</thead>
<tbody>
<tr>
<td>Control</td>
<td>14.00 ± 0.03</td>
<td>2056.18</td>
</tr>
<tr>
<td>GG_0.1%</td>
<td>3.22 ± 1.32</td>
<td>915.91</td>
</tr>
<tr>
<td>CMC_0.1%</td>
<td>1.49 ± 0.10</td>
<td>91.68</td>
</tr>
<tr>
<td>CMC_0.5%</td>
<td>0.01 ± 0.02</td>
<td>9.24</td>
</tr>
</tbody>
</table>

These results demonstrate that transport to the membrane can be controlled by the properties of the fluid under physiologically representative process conditions. As the concentration of CMC is increased, under segmentation, the mixing and flow at the membrane surface is decreased resulting in the reduction of k_{op} from 1.5 to 0.1 m/s. The Sherwood number was used to quantify the mass transfer enhancement of the system with convection compared to diffusion. Fig. 3 plots the Sherwood number (Eq. 3) against the Reynolds number. The Sherwood number increases as the Reynolds number increases for experiments when segmentation movements take place. The increase was attributed to the mass transfer enhancement from mixing by reducing the viscosity of the fluid [7].

Figure 3. Sherwood number vs Reynolds number

3.2. Microspheres and entrainment efficiency

An aqueous solution of alginate containing glucose and methylene blue was dropped into calcium chloride solutions and gelled spheres were formed instantaneously by ionotropic gelation in which intermolecular crosslinks were established between the divalent calcium ions and the negatively charged
carboxyl groups of the polymer [10,11]. The resulting beads were spherical, with diameters of mean 1.5 mm. The diameter of the beads decreased as the concentration of CaCl₂ decreased. Entrapment efficiency (EE) using alginate as a wall material was higher at low concentration of CaCl₂ (1%, w/v) (Tab. 3). Nevertheless, the EE diminished from 76% to 60% when CaCl₂ concentration was increased from 1 to 5% (w/v). These results are in agreement with those found by others authors [9]. The EE of gelled particles is usually reduced with increasing the concentration of calcium chloride in gelling medium and cross-linking time. This behavior could be explained as a denser structure was formed with less volume available for the entrapment of active substances [10]. As a result of syneresis, harder structures are formed when gelled particles are exposed to high concentration of calcium ions during extended periods of time.

Table 3. Entrapment efficiency of alginate beads.

<table>
<thead>
<tr>
<th>CaCl₂ (%)</th>
<th>Time (min)</th>
<th>EE ± std (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>5</td>
<td>5</td>
<td>76.45 ± 0.16</td>
</tr>
<tr>
<td>5</td>
<td>10</td>
<td>70.90 ± 0.33</td>
</tr>
<tr>
<td>10</td>
<td>5</td>
<td>70.45 ± 0.42</td>
</tr>
<tr>
<td>10</td>
<td>10</td>
<td>66.15 ± 0.22</td>
</tr>
</tbody>
</table>

3.3 Release of actives

Fig. 4 shows the release profile of the colorant entrapped within the alginate beads after being exposed to segmentation movements in the SIM for about 1 h. The release of the colorant and glucose was very slow reaching just 0.5% after the first hour. The entrapped colorant was suddenly released in the initial 10 minutes (about 3%) and only 0.5% more was further released in the SIM.

![Figure 4. Cumulative release of actives in the SIM with segmentation motion.](image)

This system demonstrated to have a significant effect in the delivery rates of actives. For instance, when glucose was diluted in water or other biopolymers 10% of the initial concentration was able to cross the membrane, however only 0.5% of this molecule was released after the encapsulation process. As a result very low absorption rates were obtained.

Further experiments will be done under simulated gastric and intestinal conditions studying the pH-sensitivity of the alginate beads made. Moreover other biopolymers and methods will be proved to delivery of actives.

4 CONCLUSIONS

The SIM can be used to further understand how food structure affects the physical and chemical processes of digestion. Absorption of glucose through the semi-permeable membrane increased by 20% as segmentation was applied. An increase in viscosity resulted in significant (90%) decrease of mass transfer. The ionic gelation technique proved to be an easy and efficient way to make spherical microcapsules.

ACKNOWLEDGMENTS

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REFERENCES

Appendix III

Encapsulation and Stability of Bioactive Proteins in Gel loaded Liposomes

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The entrapment efficiency and stability of liposomes loaded with alginate and a model bioactive protein alkaline phosphatase (ALP) were compared with that of conventional liposomes loaded with ALP. Multilamellar vesicles (MLV) were prepared by hydration of a thin lipid film of dipalmitoylphosphatidylcholine (DPPC) with a 20 mg/ml solution of ALP with and without 1% alginate. Once prepared the entrapped alginate was cross-linked by addition of CaCl₂ to the vesicle suspension forming liposomes with a gelled core. Entrapment of ALP was shown have 60% greater entrapment efficiency in alginate containing liposomes than liposomes prepared without alginate. Release of ALP from the gel containing liposomes at pH 2 was less than for conventional liposomes however large fluctuations in size was witnessed in the gelled liposomes when stored at pH 3.8 and at pH 7.4, possibly due to shrinking and swelling of the gel respectively. These initial findings have prompted further release and stability studies to determine the activity of the ALP following exposure to simulated gastrointestinal fluids which will reveal the potential of these particles as an oral delivery system for bioactive proteins.

Keywords: liposome, alginate, controlled release, stability, bioactive, encapsulation

1 INTRODUCTION

The incorporation of bioactive proteins in food products and delivery of such proteins in full working order to the physiological target site of the intestine can be problematic. This is due to the fact that often the functionality of bioactive proteins relies on their conformational structure which can easily be denatured by harsh processing conditions [1] and the acidic pH of gastric secretions once ingested [2]. To overcome these difficulties encapsulation strategies can be employed to stabilise and protect functional proteins. Liposomes and microspheres are just two methods used to encapsulate and deliver proteins; however there are limitations to both these delivery systems when used singularly such as stability, scale-up and undesired leaching of the entrapped molecules. The present study aims to develop and characterise a self-assembly delivery system with enteric properties by investigating the potential of loading alginate along with a modal enzyme into liposomes, then cross-linking the alginate once encapsulated. Subsequently, immobilising the protein in enteric micro-gel particles, that are coated with phospholipid bilayers.

2 MATERIALS AND METHODS

2.1 Materials

Dipalmitoylphosphatidylcholine (DPPC) was purchased from Avanti polar lipids and the sodium alginate was purchased from BDH and was high G alginate, all other reagents were purchased from Sigma-Aldrich Company (Poole, UK).

2.2 Liposome preparation

DPPC was dissolved in a 9:1 solvent mixture of chloroform and methanol. The solvent was evaporated on a rotary evaporator to obtain a dry film, which was hydrated with a solution of ALP in 100 mM Tris HCl (to give final lipid concentration of 32 μmol/ml) in a round bottomed flask at 45°C with intermittent vortexing. Unentrapped ALP was removed from the liposome suspension by repeated centrifugation (13000 rpm for 5 min) decanting the supernatant and washing the pellet with 10 mM tris HCl until no ALP was detected in the supernatant. Liposomes containing alginate were prepared using the same method however the thin lipid film was hydrated with a mixture of 1% Na alginate containing 20 mg/ml ALP. Once unentrapped APL was removed by repeated centrifugation the alginate/ALP loaded liposomes were added to 0.2M CaCl₂ for 15 min to induce gelation of the entrapped alginate. The CaCl₂ was then removed by centrifugation and the liposomes were resuspended in 10 mM Tris HCl.

2.3 Size measurements

The volume mean diameter (VMD) of the liposomes was measured using a Malvern Master Sizer 2000, (Worcestershire, UK) at 25°C, diluting 50 μl of the dispersion to the appropriate volume with double-distilled water.

2.4 Quantification of Entrapment and Release

The concentration of entrapped ALP was determined using the BCA assay. Briefly, test samples of 100 μl vesicle suspension were added
100 ml sodium dodecyl sulphate solution and 100 ml 2% sodium citrate and incubated for 1 h. Samples were then added to BCA reagent incubated at 37°C for 30 min before absorbance measurements were taken at 562 nm. Release of ALP when incubated at pH2 was also monitored, samples were taken over a 2 h period, centrifuged and the ALP content of the supernatant was quantified using the BCA assay.

2.5 Stability study

Samples of liposomes and alginate containing liposomes loaded with ALP were prepared and stored in Tris HCl maintained at pH 3.8 and 7.4 at 4°C. The liposomes were monitored for changes in size and ALP released over a period of 10 days.

3 RESULTS & DISCUSSION

The results given in Table 1 show a large difference in size between the standard MLV and MLV containing alginate. This increase in size is likely to be due to aggregation where residual alginate is present. Entrapment was also seen to increase in the vesicles containing alginate. This is thought to be a consequence of immobilisation of the ALP within the alginate gel network ultimately reducing leakage through the liposome bilayer. Multilamellar vesicles are expected to be more stable than other liposome types due to the many bilayers surrounding the encapsulated material [2]. However the addition of alginate appears to increase the stability releasing 7% less ALP when incubated for 2 hours at 37°C at pH2. This stability was highlighted further by evaluating release over 48 h at pH 2 which revealed the alginate containing liposomes releasing 70% ALP compared with 32% retained by the standard liposomes (results not shown).

<table>
<thead>
<tr>
<th>Sample</th>
<th>Mean Diameter (μm)</th>
<th>% Entrapment of ALP</th>
<th>% Released in 2h at pH 2</th>
</tr>
</thead>
<tbody>
<tr>
<td>Liposomes</td>
<td>30.8 ± 3.1</td>
<td>0.9 ± 0.3</td>
<td>19.6 ± 1.4</td>
</tr>
<tr>
<td>Gelled liposomes</td>
<td>13.4 ± 0.1</td>
<td>1.6 ± 0.3</td>
<td>13.8 ± 1.3</td>
</tr>
</tbody>
</table>

Table 1: Liposomes and liposomes containing alginate gel. Comparison of particle size, % entrapment of ALP and release following 2 h at pH 2. All measurements were obtained for at least three independently synthesized batches and Mean ± S.D. is reported for each sample.

The effects of pH on the stability of both vesicle types was additionally analysed by monitoring size over a period of 10 days. Fig.2 shows samples stored at pH 3.8. The alginate containing vesicles reduced in size, possibly a result of the alginate gel contracting. The standard MLV revealed a slight increase in size when stored at pH 3.8 which is a possible indication instability. When this is compared with MLV stored at pH 7.4 (Fig.3) the particle size remains constant for 10 days. Interestingly, the alginate containing liposomes increased from 30 μm to 45 μm after 10 days at pH 7.4. This may be due to the alginate gel gradually swelling. Further studies are in progress monitoring release of ALP from the particles at pH 7.4 and the activity of the encapsulated ALP following exposure to simulated gastric fluid.

4 CONCLUSION

Liposomes loaded with alginate that is then gelled by addition of CaCl2 may be a potential approach for the encapsulation of bioactive proteins, to protect them from harsh processing environments and acidic pH of the stomach once ingested.

References