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Mesoscale modelling of polymer aggregate digestion

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1. Introduction

Food digestion is a complex cascade of chemical and physical processes spanning multiple length scales (Bornhorst et al., 2016; Wang et al., 2010), ranging from centimetric dimensions for food ingested through the mouth down to the molecular scale for nutrients broken down and absorbed in the stomach and the intestines. As studies have established links between dietary habits and many common health issues, e.g. obesity and diabetes (Bornhorst et al., 2016) understanding digestion is of broad biomedical relevance. Specifically, a better insight into the process would facilitate the design of so-called ‘functional foods’ that aside from nutritional benefits are associated with an improved state of health and/or reduction of the risk of some diseases (Vivada-Martos et al., 2010; Ozen et al., 2012). The effect of functional and other foods on health is to a large extent controlled by their rate of digestion; for example, resistant starches, i.e., ones that are not broken down until they are transported to the large intestine, are known to have beneficial health effects (Bird et al., 2009).

Food digestion starts with mastication in the mouth. There, food is broken down mechanically, lubricated by saliva and converted into a cohesive mass known as a bolus (Singh et al., 2015; Bornhorst and Singh, 2012) that consists of particles of typical size of ~ 1 mm (Jalabert-Malbos et al., 2007; Bornhorst and Singh, 2012). The bolus is then transported to the stomach where it is broken down chemically through hydrolysis by the hydrochloric acid and the enzymes in the gastric juices, as well as mechanically via muscular contractions; limited nutrient adsorption also occurs in the stomach (Singh et al., 2015). After gastric sieving, which only allows particles of size smaller than ~ 1 mm to pass (Schulze, 2006), the mixture of partially broken down food and secretions from the digestive tract (chyme) is then transported to the small intestine where the pH is neutral to basic (Boland, 2016). The small intestine is the site of further enzymatic and mechanical breakdown and much of the absorption of nutrients occurs there (Cleary et al., 2015). Finally, a low-viscosity slurry reaches the large intestine where the key processes are microbial breakdown and water absorption (Cleary et al., 2015).

Given the complicated and multiscale nature of the digestive pathway, and the varied structure of the biopolymeric molecules relevant to the human diet, formulating models of digestion is very demanding and progress is only likely to be made by using a range of methods relevant to different length scales. Much of the existing work is based on continuum approaches, see e.g. Taghipoor et al. (2012), Moxon et al. (2016) and the reviews in Cleary et al. (2015), Ferrua et al. (2011), and...
Ferrua and Singh (2015). In particular, some authors have formulated anatomically accurate 3D computational models of food breakdown in the mouth (Cleary et al., 2015) and of gastric digestion (Cleary et al., 2015; Ferrua and Singh, 2010, 2015; Ferrua et al., 2011). Other authors have also worked towards developing in vitro models for the experimental study of food digestion, e.g. the gastric simulator described by Kong and Singh (2008, 2010). Although some effort has been dedicated to studying digestion at the mesoscale, for example via the lattice Boltzmann method (Wang et al., 2010), to the best of our knowledge, the approach described of this article, using a mesoscale algorithm to study the dynamics and stability of the polymeric aggregates that arise in the intermediate stages of the breakdown of complex carbohydrates, has not yet been explored. The method we use explicitly models the flow of polymer-solvent mixtures, eliminating the need for approximate constitutive relations and the corresponding assumptions. However, feasible simulations do require coarse-graining molecular detail.

We focus on the dynamics of polymeric aggregates in a viscous medium. Although we do not aim to mimic physiological conditions exactly, our results are most pertinent to food digestion in the stomatch. Rather than attempting to simulate a specific biopolymer or to describe the complex mixture of biomolecules encountered in the digestive tract, we use a simple model that allows insight into some of the generic mechanisms controlling digestion. Our polymeric aggregates, which we refer to as boluses, are initially approximately spherical in shape and consist of monodisperse linear bead-spring chains with no bonding rigidity, and we assume that a fraction of the beads can form physical cross-links. (Note that we use the term ‘bolus’ to refer to a generic aggregated mass rather than in its specific sense of a lubricated conglomerate of typically millimeter food particles formed during mastication (Jalabert-Malbos et al., 2007; Bornhorst and Singh, 2012).

We first consider the dynamics of such boluses in a quiescent fluid, determining the key parameters that control whether they are stable with respect to diffusion. We next study how two of the major factors at play in the digestive tract, chemical breakdown of the polymers and shear flow, affect bolus evolution. Finally, we investigate the combined action of simple shear and chemical breakdown of the polymers. The predictions of our model, while not directly comparable to a specific system, are relevant to the digestion of various biopolymers, e.g. starch, which is a major energy source in the typical human diet (Shrestha et al., 2015; Liu et al., 2017).

2. Methods

2.1. Polymer model

We consider a coarse-grained, bead-spring model of monodisperse linear polymer chains in which the bonds are approximated by harmonic springs, yielding the potential (Allen and Tildesley, 1987):

$$u_{\text{bond}} = \frac{1}{2} k_{\text{bond}} \sum_{i=1}^{N_{\text{bond}}} (|\Delta r_i| - l_0)^2$$

where $r_i$ is the position vector of bead $i$, $N_{\text{bond}}$ is the number of beads in a single chain, the resting bond length is $l_0 = \sigma$, $\sigma$ being the bead diameter, $k_{\text{bond}} = 10^9 k_B T$, where $k_B$ is the Boltzmann constant and $T$ is the temperature.

We model the inter- and intra-chain interactions between beads with the truncated Lennard-Jones potential (Allen and Tildesley, 1987):

$$u_{\text{LJ}} = \begin{cases} 
\epsilon + 4\epsilon \left( \frac{\sigma}{r} \right)^{12} - \left( \frac{\sigma}{r} \right)^6 & r \leq r_{\text{cut-off}} \\
0 & r > r_{\text{cut-off}}
\end{cases}$$

where $r$ is the distance between the centres of the beads and $\epsilon$, which is measured in units of $k_B T$, characterizes the interaction strength. For beads that do not form cross-links, we use the Weeks-Chandler-Andersen potential (Weeks et al., 1971), which we obtain from Eq. (2) by setting the cut-off radius to $\sigma$ and $\epsilon' = \epsilon / 2^{1/6}$. This choice of interaction potential implies that we are modelling a good solvent (Ryder and Yeomans, 2006).

To model physical cross-linking, which can act both within and between polymer chains, we modify the Lennard-Jones potential so that interactions between individual polymer beads can be attractive. Upon generating the initial conditions for each simulation, there is a probability $p_{\text{link}}$ that each bead will be able to form links. We model the force between two linking beads with an offset Lennard-Jones potential,

$$u_{\text{link}} = u_{\text{LJ}} + u_{\text{offset}}$$

where we choose the offset so that the force acting between the two beads, $F = -\nabla u_{\text{link}}$, vanishes at $r = r_{\text{cut-off}}$:

$$u_{\text{offset}} = \begin{cases} 
\frac{\sigma^2}{2 r_{\text{cut-off}}^2} \lim_{\Delta r \to 0} \frac{\partial u_{\text{LJ}}}{\partial r} & r \leq r_{\text{cut-off}} \\
0 & r > r_{\text{cut-off}}
\end{cases}$$

For linking beads, $\sigma' = \sigma$ and $r_{\text{cut-off}} = (5/2)\sigma$ in both $u_{\text{LJ}}$ and $u_{\text{offset}}$. $u_{\text{link}}$ has a minimum of $\approx 10^{-2} \epsilon$ at approximately the same $r$ as $u_{\text{LJ}}$, $r_{\text{min}} \approx 7/4 \sigma$. Offsetting the interaction potential in this way is standard in MD simulations, and a discussion of using an offset linear in $r$ can be found in Section 5.2.4 in Allen and Tildesley (1987). Note that the physical cross-links, which act both within and between polymer molecules, are much weaker than the intramolecular chemical bonds holding the chains together. For comparison, if we take the Taylor series of $u_{\text{link}}$ about $r_{\text{ens}}$ and truncate it to second order, we obtain an effective spring constant of $k_{\text{eff}} = u''(r_{\text{ens}}) \approx 56.8 \epsilon / \sigma$, with $\epsilon \approx k_B T$, whereas $k_{\text{bond}} = 10^9 k_B T$.

In order to prevent the repulsive forces from diverging during initialization we normalize the forces arising from the WCA potential and $u_{\text{link}}$ in the following way: rewriting the force as $F = -\nabla u = \nabla F_0$, with $r$ being the vector connecting the centres of the two beads, for $|F_0| > F_{\text{max}} = 10^5 k_B T / \sigma$, we use the expression $F = r F_{\text{max}} F_0 / |F_0|$. The force normalization is only relevant if two beads overlap significantly. Our algorithm for bead initialization allows such overlaps, but they are quickly eliminated as the aggregates evolve and unlikely to occur elsewhere in our simulations as the bead-to-bead interaction potential becomes strongly repulsive at short distances.

We generate the initial positions of the beads so that they form an aggregate (bolus) of approximately spherical shape with all beads within a sphere of radius $R_{\text{sphere}} = 6 \sigma$ whose centre coincides with that of the simulation box. We specify the desired polymer volume fraction $\rho$ in the sphere and generate $N_{\text{poly}}$ polymer chains such that $\rho$ is not exceeded, $N_{\text{poly}} = 6\pi V_{\text{sphere}} / (\pi N_{\text{bead}} \sigma^2)$, where the floor function $x$ acting on the real number $x$ returns the largest integer $\leq x$, see Hase-winkel (2002). For each set of parameters, we perform $N_{\text{ens}} = 20$ simulations with different random initial conditions and average the results over the ensemble.

The process of digestion involves enzymes that hydrolyse the chemical bonds between monomers and thus break down the polymeric chains. As a simple way of modelling the enzymatic hydrolysis, we introduce random bond cleavage to the model. We do this by introducing a rate constant $k_{\text{break}}$, which characterizes the probability that the bond between any two beads is broken at each streaming step; if a bond is cleaved, the two beads participating in it are no longer connected via the harmonic potential (1). All bonds are equally susceptible to attack: this mode of enzyme degradation of carbohydrates is known
2.2. Multi-particle collision dynamics

We simulate the flow via multiparticle collision dynamics (MPCD), a mesoscale technique for solving the Navier-Stokes equations that treats the solvent as a collection of point particles which move ballistically during streaming steps and exchange momentum in collision steps (Yeomans, 2006). The MPCD implementation used here follows the work of Zöttl and Yeomans (Zöttl and Yeomans, 2019a, b), see references therein, in particular Gompper et al. (2009). The main difference with the approach from Zöttl and Yeomans (2019a, b) is that here we also include physical cross-linking. In a streaming step of duration $\delta t$, a fluid particle changes its position $\mathbf{r}_i$ according to

$$r_i(t + \delta t) = r_i(t) + v_i(t)\delta t$$

where $v_i$ is the particle velocity. Between streaming steps, the particles are sorted in cubic cells of length $L_0 = \sigma$ and exchange momentum in collision steps according to a collision rule that employs an Andersen thermostat (Zöttl, 2014):

$$v_i(t + \delta t) = v_{cell}(t) + v_{rand}(t) + v_p(t) + v_e(t)$$

where $v_{cell}$ is the centre-of-mass velocity of the cell, $v_{rand}$ is a random velocity obeying the Maxwell-Boltzmann distribution at temperature $T$, and the terms $v_p$ and $v_e$ ensure that linear and angular momentum are conserved (Gompper et al., 2009). The parameters for the MPCD fluid are the same as in Zöttl and Yeomans (2019b) and correspond to a viscous flow with a low Reynolds number: the number density of the fluid particles is $N_{poly} = 10/\sigma^3$, the time step is $\delta t = 0.02/\sqrt{\sigma^2/(k_B T)}$, being the mass of an individual particle. The quantity $\tau_0 = \sqrt{m^2/(k_B T)}$ has the dimension of time, and throughout the text, it is the implied time unit wherever no other is specified. The implied units of length, mass and energy are respectively $\sigma$, $m$ and $k_B T$.

During the streaming step, the forces acting on the polymer beads are calculated from Eqs. (1)–(3) and their positions are computed via the velocity Verlet algorithm (Allen and Tildesley, 1987) at intervals of $\delta t_{poly} = \delta t/50$. The polymer beads have mass $m_p = 10m$ and are included in the collision step (Yeomans, 2006; Zöttl and Yeomans, 2019b). This two-way coupling between the solvent and the polymer molecules means that the swelling of the aggregate is naturally accounted for. MPCD has been used to study related problems such as the dynamics of the transition from a swollen to a collapsed state in a dilute polymer solution, see e.g. Kikuchi et al. (2002, 2005). Other ways of using MPCD to model polymeric solution have been proposed in the literature, e.g. the two approaches described by Ruiz-Franco et al. (2019), both of which generate coarser graining based on monomer density profiles for simulations with explicitly modelled monomers.

We study the temporal evolution of polymeric aggregates in an unbounded, initially quiescent fluid and in shear flow. To this end, we employ a cubic simulation box of size $L = 48\sigma$, and simulate a time period of $t_{sim} = 10^8/\delta t$. For simulations involving an unbounded fluid, we apply periodic boundary conditions to all walls. To simulate shear flow, we introduce planar solid walls that are situated at $x = \pm L/2$ and move with a velocity $v_x = \pm v_{wall}$. This corresponds to an approximately linear $v_x(x)$ with a shear rate $\dot{\gamma} = 2v_{wall}/L$. In simulations that involve solid walls, the walls contain virtual particles that interact with those of the fluid and polymer according to a bounce-back rule (Zöttl, 2014).

### 3. Results

In the simulations we discuss below, we study the evolution of aggregates comprised of fully flexible polymer chains with a Kuhn length of $\sim \sigma$ comprised of $N_{bead} = 20$ coarse-grained beads. We explore the effect of various parameters, namely, the average fraction of beads that can form links ($p_{link}$), the interaction energy parameter ($\epsilon$), the rate of enzymatic hydrolysis ($k_{break}$) and the shear rate of imposed shear flows ($\dot{\gamma}$). We summarize the key parameters in Table 1.

For all quantities of interest $X(t)$, we calculate the ensemble mean value,

$$\langle X(t) \rangle = \frac{1}{N_{poly}} \sum_{i=1}^{N_{poly}} X_i(t),$$

where $k$ labels a given set of initial conditions. The position vector of the bolus centre of mass for the simulation with the set of initial conditions labelled $k$ is

$$r_{cm} k(t) = \frac{1}{N_{bead}} \sum_{j=1}^{N_{bead}} r_{CM} \cdot \mathbf{e}(t)$$

A useful quantity to characterize the shape and dimensions of the irregularly-shaped boluses is the gyration tensor. Its diagonal components are defined as

$$G_{ii} k(t) = \frac{1}{N_{bead}} \sum_{j=1}^{N_{bead}} (r_{bead}(t) - r_{cm} k(t)) \cdot \mathbf{e}(t)$$

where $N_{bead} = N_{poly} N_{bead}$ is the total number of beads in the bolus and $\mathbf{e}(t)$ is a Cartesian basis vector. Other authors have used analogous definitions of $G_k$ to characterize individual polymer molecules, see e.g. Mattice and Suter (1994) and Liebetreu et al. (2018). In particular, gyration tensors have been defined for individual ring polymers (Liebetreu et al., 2018; Chen et al., 2013) and single-chain polymeric nanoparticles (Fornacon and Moreno, 2018).

The bolus gyration radius is related to the trace of the gyration tensor $G_k$ as

$$R_g^{2, bol} k = \text{tr}(G_k)$$

for polymers that do not form cross-links, the individual molecules diffuse away and the aggregate disperses (Fig. 1), leading to a $\langle R_g^{2, bol}(t) \rangle$ which is linear in $t$ at long times. In contrast, for boluses with a sufficiently high number of strong cross-links, $\langle R_g^{2, bol} \rangle$ approaches a stationary value. For this reason, the quantity $s = \dot{\gamma}/(R_g^{2, bol}(t))/R_g^{2, bol}(0)$ which characterizes the rate of expansion of the bolus surface, is useful in describing the different modes of bolus behaviour.

Bolus evolution at short times is dominated by repulsive forces due to overlapping beads in the initial condition, leading to a sharp maximum of $\langle R_g^{2, bol}(t) \rangle$ at $t = 0$, see e.g. Fig. 2. This repulsive Lennard-Jones interaction is short-ranged and after going through this maximum in size, the bolus contracts due to attractive forces, typically going through a shallow minimum in $\langle R_g^{2, bol}(t) \rangle$. At still longer time scales, at $t \geq t_0 = t_{sim}/20 = 100\tau_0$, $\langle R_g^{2, bol}(t) \rangle$ is determined by the balance between

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Significance</th>
<th>Studied range</th>
</tr>
</thead>
<tbody>
<tr>
<td>$p_{link}$</td>
<td>probability that every individual bead is initialized as a linking bead</td>
<td>0 – 1</td>
</tr>
<tr>
<td>$\epsilon$</td>
<td>interaction strength, see eqs. 2-4</td>
<td>1 – 10k_BT</td>
</tr>
<tr>
<td>$k_{break}$</td>
<td>probability that a bond is broken in a given time step</td>
<td>0 – 10^{-3}</td>
</tr>
<tr>
<td>$W_i$</td>
<td>dimensionless shear rate, defined as the product of the dimensional shear rate and the bolus characteristic time, $t_{bolus}$</td>
<td>0 – 58</td>
</tr>
</tbody>
</table>
cross-linking interactions holding the bolus together and the diffusion of the polymers driving the dispersion of the aggregate. As this is the regime we are interested in, we focus on $t > t_0$.

3.1. Boluses in a quiescent fluid

We start by investigating the stability of boluses in a quiescent fluid and its dependence on the properties of the constituent polymers.

3.1.1. Varying the linking probability $p_{\text{link}}$

In Fig. 1, we see snapshots of boluses at the beginning and the end of simulations. The polymeric molecules comprising the aggregates in Fig. 1(a) can form no physical cross-links, whereas the fraction of linking beads in Fig. 1(b) is $\approx 40\%$. As the comparison between these two cases indicates, introducing cross-linking interactions qualitatively changes the behaviour of polymeric aggregates, causing them to approach a stationary radius rather than disperse over time. The plots of the ensemble-averaged squared gyration radius $\langle R^2_{\text{bol}}(t) \rangle / \langle R^2_{\text{bol}}(0) \rangle$ in Fig. 2 demonstrate that as $p_{\text{link}}$ is increased from 0, bolus behaviour transitions from dispersing over time to attaining a stationary gyration radius (see also Supplementary Movies 1 and 2). Looking at the bolus expansion rate $s$ shown in the inset of Fig. 2 suggests that this is quite a sharp transition at $p_{\text{link}} \approx 0.3$. This value is related to the fraction of linking beads required to bind all molecules in the aggregate together via cross-links and is therefore related to a percolation threshold above which the entire aggregate is bound by cross-links.

However, this is not a simple geometric percolation transition because it relates to the formation of a network of cross-links within a
finite aggregate instead of an infinite medium. Moreover, the stability of the bolus is controlled not only by the number of cross-links in it but also by their strength, as well as additional factors, such as the initial volume fraction of polymer and the length of the polymeric chains. We discuss the most pertinent of these parameters below and choose the ensemble of boluses with \( p_{\text{link}} = 8/N_{\text{bead}} \cdot N_{\text{bead}} = 20, \rho = 0.4 \) and \( \varepsilon = 5k_B T \) as a reference system since these parameters yield aggregates that are stable in quiescent conditions.

Note that even at \( p_{\text{link}} = 1 \), the bolus radius varies with time. The initial bead positions are random, which means that beads that are within the range of the Lennard-Jones potential from each other can come closer or spread apart as time progresses, depending on whether the initial distance between the beads is greater or smaller than the equilibrium value. Furthermore, the energy of physical cross-linking is of the same order of magnitude as the thermal energy of the beads, meaning that the polymer configurations are perturbed by thermal fluctuations, which are naturally included in the MPCD method.

3.1.2. Varying the interaction energy parameter \( \varepsilon \)

In Fig. 3 we show the long-time behaviour of the squared bolus gyration radius \( \langle R_g^2 \rangle \) for different values of the interaction energy \( \varepsilon \). The figure demonstrates that merely having a \( p_{\text{link}} \) that ensures cross-links span the entire aggregate is not sufficient to make the bolus stable with respect to diffusion and that \( \varepsilon \) also needs to be above a threshold value for the aggregates to remain intact at long times. For our model system \( (N_{\text{bead}} = 20, \rho = 0.4, p_{\text{link}} = 4.0/N_{\text{bead}}) \), we see that the long-time bolus expansion rate drops sharply at \( \varepsilon \approx 3k_B T \).

3.1.3. Chemical breakdown

We study the effect of polymer hydrolysis by varying the rate constant \( k_{\text{break}} \) over several orders of magnitude. We choose the values of \( k_{\text{break}} \) so that the number of bonds broken during the course of the simulations \( t_{\text{sim}} \) varies between 0 and the total number of bonds in the bolus. As seen in Fig. 4, which contains plots of the bolus ensemble-averaged squared gyration radius \( \langle R_g^2 \rangle \) at various \( k_{\text{break}} \), chemical breakdown in the model drives boluses that are stable in the absence of hydrolysis to disperse if \( k_{\text{break}} \geq 10^{-6} \), corresponding to the cleavage of \( \geq 10\% \) of the bonds in the aggregate over the course of the simulation.

The bolus evolution in the case of high \( k_{\text{break}} \) has an unexpected feature: in this case complete hydrolysis of the polymers occurs at \( t \ll t_{\text{sim}} \) and the bolus is reduced to a collection of individual beads. This allows the cross-linking beads to interact more strongly, forming a greater number of cross-
3.2. Tumbling, tank-treading and breakdown in shear flow

Boluses under shear exhibit three different regimes depending on the imposed shear rate $\dot{\gamma}$. For boluses that are stable under quiescent conditions, we nondimensionalize $\dot{\gamma}$ with a characteristic time ($t_{\text{bolus}}$), which we define as the time required for $\langle R_g^2 \rangle_{\text{bolus}}$ of a bolus with identical parameters to reach 95% of $\langle R_g^2 \rangle_{\text{bolus}}$ in the absence of shear. Note that, as before, we disregard the initial period in which $\langle R_g^2 \rangle_{\text{bolus}}$ goes through a maximum. For this reason, when determining ($t_{\text{bolus}}$), we start measuring $\langle R_g^2 \rangle_{\text{bolus}}(t)$ at $t = t_{\text{lim}}/20$. The dimensionless bolus Weissenberg number is then $W_i = \dot{\gamma} t_{\text{bolus}}$, with ($t_{\text{bolus}}$) > $t_{\text{lim}}$; for our model system ($N_{\text{bead}} = 20, \rho = 0.4, p_{\text{link}} = 4.0/N_{\text{bead}}, t_{\text{wall}} = 0, k_{\text{break}} = 0$), ($t_{\text{bolus}}$) = 140$\tau_0$.

We now discuss the different regimes of bolus behaviour in shear flow. i) At $W_i \ll 1$, the bolus is not significantly perturbed by the shear flow and both $\langle R_g^2 \rangle_{\text{bolus}}$ and $s$ are unchanged with respect to the quiescent case (Fig. 5(a)). ii) At intermediate $W_i$, $W_i \sim 1$, boluses are deformed and move collectively in the direction of the flow (Fig. 5(b), Supplementary Movie 4). Individual polymer chains break off from some of the boluses in an ensemble (Supplementary Movie 5), but overall, the bolus maintains its structure. The transition between regimes i) and ii) occurs when the energy dissipated due to viscous friction over the characteristic time for the bolus ($t_{\text{bolus}}$) becomes much larger than the

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total energy of the cross-links in it. iii) At $Wi \gg 1$, the boluses are broken apart by the shear flow (Fig. 5(c), Supplementary Movie 6).

The final bolus size $\langle R_{g \text{ bol}}(t) \rangle$ for $Wi \gg 1$ is orders of magnitude greater than the box length $L$. This means that the box no longer adequately simulates an infinite medium, that intermolecular interactions are stronger than they would be in an infinite box, and that the results we report for this case at long times are approximate; the same is true of simulations in which individual molecules are separated from the bolus by the shear flow. However, we expect that the finite size of the box does not introduce a significant error in the quantities of interest to us in the latter case because the splitting of individual molecules from an aggregate of ~30 chains has a minor effect on its gyration radius.

In regime ii), in which the aggregates move approximately as solid bodies, there are two limiting modes of motion that the bolus can exhibit: tumbling and tank-treading. In the first of these, tumbling, the polymeric chains experience large conformational changes and alternate between stretched and collapsed states. In the second one, tank-treading, individual beads rotate about the bolus centre of mass and the conformation of the chains is approximately constant. See Chen et al. (2013) and Formanek and Moreno (2018) for discussion of these modes for individual ring polymers and nanoparticles consisting of a single polymeric chain, respectively.

We use two correlation functions to describe this collective motion, on the scale of the entire bolus, at intermediate shear rates. The first one is the cross-correlation function of the diagonal components of $G$ in the flow and gradient direction, which characterizes tumbling,

$$C_{xx} = \frac{\langle \delta G_{xx}(t) \delta G_{xx}(0) \rangle}{\sigma_{G_{xx}(t)} \sigma_{G_{xx}(0)}}$$

where

$$\delta G_{xx}(t) = G_{xx}(t) - \langle G_{xx}(t) \rangle$$

and

$$\sigma_{G_{xx}(t)} = \sqrt{\frac{N_{\text{ens}}}{N_{\text{ens}} - 1} \left( \langle G_{xx}(t)^2 \rangle - \langle G_{xx}(t) \rangle^2 \right)}$$

are the standard deviations of the diagonal components of the gyration tensor. We choose the offset time $t_0 = t_{\text{sim}}/20$ so that the repulsion-dominated initial period during which the bolus gyration radius goes...
Fig. 5. Snapshots of the evolution of boluses with \( N_{\text{bend}} = 20, \rho = 0.4, \rho_{\text{link}} = 8/N_{\text{bend}} = 20, \varepsilon = 5.0 k_B T \) subjected to flows with different shear rates. The shear is created by the motion of the horizontal solid walls as indicated by the arrows in the first frame in a. The scale bars are of length 10\( \sigma \).

- **Regime i)** - at low Wi (Wi = 0.58), the bolus is not significantly affected by the flow.
- **Regime ii)** - at intermediate Wi (Wi = 5.8), the bolus performs a tumbling/tank-treading motion.
- **Regime iii)** - at high Wi (Wi = 29), the bolus is broken apart by the flow.
through a maximum does not contribute to the correlation functions.

Negative peaks in the cross-correlation function $C_{ax}$ are a hallmark of tumbling motion (Formanek and Moreno, 2018). These peaks arise because the polymer chains are preferentially stretched along the flow direction, but thermal fluctuations cause stretching in the gradient direction. This causes the chains to contract and subsequently extend along the flow.

The correlation function that characterizes tank-treading is

$$C_{\text{angle}}(t) = \frac{\langle \sin(2\beta(t_0))\sin(2\beta(t)) \rangle}{\langle \sin^2(2\beta(t_0)) \rangle} = \frac{\sum_{k=1}^{N_{\text{bol}}} \sum_{i=1}^{N_{\text{dis}}} \sin(2\beta_{ik}(t_0)) \sin(2\beta_{ik}(t))}{\sum_{k=1}^{N_{\text{bol}}} \sum_{i=1}^{N_{\text{dis}}} \sin^2(2\beta_{ik}(t_0))}, \tag{14}$$

where $\beta_{ik}$ is the angle between the vector connecting the bead with position vector $r_{ik}$ to the bolus centre and the instantaneous first principal component of the bolus bead positions. The principal components of the instantaneous set of bead positions are basis vectors defined through a linear transformation of the Cartesian basis in which we record the positions. The transformation is defined such that the principal components maximize variance and are orthogonal to each other (Jolliffe, 2002). The principal components correspond to the axes of an ellipsoid fit to the distribution of bead positions, and the first principal component corresponds to this ellipsoid’s principal axis along which statistical variation is greatest Jolliffe (2002), thus defining the main bolus axis. $\beta_{ik}$ is therefore defined through

$$\cos(\beta_{ik}(t)) = \frac{\langle r_{ik} - r_{x_k}(t) \rangle \cdot \hat{e}_k}{|r_{ik} - r_{x_k}(t)|}, \tag{15}$$

where $\hat{e}_k$ is a unit vector in the direction of the first principal component.

Damped oscillations of $C_{\text{angle}}$ with time are a characteristic sign of tank-treading motion (Chen et al., 2013; Formanek and Moreno, 2018). In contrast with tumbling, which causes polymer molecules to alternate between stretched and compressed conformations, tank-treading motion occurs with rotation of the individual beads around the bolus centre of mass that maintains the conformation of individual molecules approximately constant (Formanek and Moreno, 2018). This periodic stretching and contraction of individual molecules in the direction of the flow is reflected at the scale of the entire aggregate through the components of the gyration tensor.

The correlation functions $C_{ax}$ and $C_{\text{angle}}$ are plotted in Fig. 6(a) and (b). They indicate that, as $Wi$ is increased from 0, the boluses studied here first exhibit slow tumbling motion at $Wi \sim 1$ (thick solid curve, Fig. 6(a)), and then move in a way that combines tumbling and tank-treading (dotted and dashed curves, Fig. 6) at higher $Wi$.

3.2.2. Synergy of shear flow and chemical breakdown

Finally, we consider the case of bolus evolution in the presence of both shear flow and chemical breakdown, which is particularly relevant to modelling the digestive tract where muscle contractions induce mixing and enzymes catalyse hydrolytic reactions. Fig. 7, which contains data on $(R_{g, \text{bol}}(t_{\text{break}}))$ at different dimensionless shear rates, illustrates the synergistic effect of the two factors. The graph demonstrates that the combination of fast polymer hydrolysis and a low Weissenberg number is more efficient than either of the two on its own (see also Supplementary Movie 7).

Fig. 7 shows that flows with $Wi = 0.29$, which in the absence of chemical breakdown have a negligible effect on bolus size, lead to a considerable increase in the squared gyration radius at long times and $k_{\text{break}}$ high enough to cleave a substantial fraction of the chemical bonds in the aggregate ($(R_{g, \text{bol}}^2)(t_{\text{break}})$ increases by $\approx 30\%$ at $k_{\text{break}} = 10^{-5}$ and approximately twofold at $k_{\text{break}} = 10^{-4} - 10^{-5}$).
pigs shows a particle distribution with areas ranging from 10$^{3}$ to 10$^{7}$ s$^{-1}$ for amylose and ~10$^{3}$ to ~10$^{9}$ s$^{-1}$ for xanthan. Shear rates in the digestive tract have not been accurately measured (Dikeman and Fahey, 2006) but based on simulation data (Ferrua et al., 2011), we can estimate them to be ~1 s$^{-1}$. Our estimates predict that the shear rates required to perturb the aggregates formed from amylose and xanthan are much higher than this value, implying that mixing in the digestive tract would play no role in the breaking down of boluses with such small dimensions (0.1 µm and ~1 µm, respectively).

Our model can also serve as a first approximation to describing protein gels and their digestion by proteinases if we disregard the differences in properties between the amino acid residues that comprise the protein molecules, as well as features of protein spatial structure such as α-helices. Even though proteins are typically charged, the comparatively high ionic strength in the digestive tract (~0.1 mol/L, see Lindahl et al. (1997) leads to a Debye length of ~1 nm, meaning that electrostatic interactions are effectively screened and can be neglected for the purposes of a coarse-grained model.

An alternative way of interpreting the model is to equate the initial aggregate size, $R_{\text{sphere}} = \theta$, to the approximate radius of the smallest particles in digesta observed by Bornhorst et al. (2013), $R_{\text{sphere}} \approx 32 \mu$m. This is equivalent to assuming that each polymer in the bolus represents multiple entangled polymeric chains. Taking the linear density of amylose, $\rho = 2.39 \times 10^{-6} \text{kg.m}^{-1}$ (Stokke et al., 1987), we find that the range of shear rates we explore falls between 5 and 5 $\times$ 10$^3$ s$^{-1}$, and that the bead diameter is $\sigma = 5.3 \mu$m. In this mapping, given the presence of hydrolysis, a physiologically relevant shear rate of $\dot{\gamma} \approx 11 s^{-1}$ (Wi = 0.15) is sufficient to significantly affect the digestion of boluses.

The coarse-grained mesoscale model discussed in this paper provides insight into generic mechanisms of polymer aggregate breakdown relevant to modelling the process of digestion. Our work considers some of the main factors at play in the digestive tract, but there are many simplifying assumptions that require further investigation. The physical cross-links present in the model can serve as an approximation of the hydrogen bonds (Liang and Ludescher, 2015) and other non-covalent interactions that act between starch molecules. However, native starch consists of branched-chain amylpectin and linear amyllose molecules (Buleon et al., 1998), which can be organized in complex structures known as granules. Granules vary between 1 and ~100 µm in size (Buleon et al., 1998) and contain alternating amorphous and crystalline layers, although these are typically disrupted during food processing (Zhang et al., 2014) and further perturbed during digestion. First steps towards modelling more realistic polymer architectures could be to consider chain branching, polydispersity or the effects of electrostatic interactions.

In our model for polymer hydrolysis the chemical bonds break down spontaneously at a constant rate regardless of their position within the bolus. An enzyme that hydrolyses amylose according to this mechanism is β-amylase, which is found in plants, see e.g. Bird and Hopkins (1954). A more realistic model of hydrolysis due to the α-amylase present in human saliva might follow a multiple attack mechanism in which the enzyme at first binds to a random site along the carbohydrate chain and then hydrolyses several bonds before detaching from it (Robyt and French, 1967). Moreover, being macromolecules themselves, enzymes diffuse through the polymeric aggregates at a finite rate and a more detailed model should account for enzyme diffusion. In this case, chains at the surface of the aggregate would be attacked first; individual linking beads would have more time to diffuse after hydrolysis and would be less likely to form a dense core as they do in our simulations (see the snapshot for high $k_{\text{break}}$ in Fig. 4).

It would be very interesting to compare the simulation results to experiments on model systems: fully realistic numerical models of digestive processes are currently out of reach, but developing simpler numerical and experimental model systems in tandem will help to identify and understand the most important physical and chemical processes which contribute to digestion.
5. Conclusions

Our simulations of the dynamics of physically cross-linked aggregates consisting of linear polymers (boluses) provide insight into the process of digestion at the mesoscale. The coarse-grained models we employ allow us to pinpoint the key factors controlling the breakdown of such aggregates. We demonstrate that in a quiescent fluid, the stability of polymeric boluses with respect to diffusion is mainly controlled by the fraction of individual beads within them that can form physical cross-links \( \left( p_{\text{link}} \right) \) and the energy of the cross-linking interactions \( \varepsilon \). The bolus surface expansion rate \( s \) sharply decreases to zero at a value of \( p_{\text{link}} \) that corresponds to a network of cross-links encompassing the entire aggregate provided that the cross-links are strong enough to hold the molecules together, i.e., that \( \varepsilon \) is above a threshold value.

Two major factors control the breakdown of biopolymers in the digestive tract - enzymatic hydrolysis and shear flow. Our simple model of enzymatic polymer hydrolysis posits that all chemical bonds within the bolus degrade at random with the same probability. It demonstrates that a rate of hydrolysis which cleaves a significant fraction of the bonds within an aggregate over the simulated time period causes boluses that are stable in the absence of hydrolysis to disperse.

We also investigate the effect of simple shear on polymeric aggregates by introducing two horizontal solid walls moving in opposite directions. We observe that aggregates exhibit three different regimes depending on the magnitude of the Weissenberg number \( Wi \) for the flow: i) At \( Wi \ll 1 \), the flow does not have an appreciable effect on the aggregates. ii) At \( Wi \sim 1 \), boluses move collectively along the flow in a regime that combines tumbling and tank-treading, and individual molecules may detach from the main aggregates. iii) At \( Wi \gg 1 \), the shear flow is sufficiently strong to break the aggregates down completely.

Finally, we show that when combined, shear flow and chemical breakdown of the polymers act in synergy to disperse the aggregates. In the limit of fast chemical breakdown in which the polymeric chains are completely split into individual beads over the duration of the simulation, flows with \( Wi \sim 0.1 \) significantly aid bolus dispersion. This is in contrast to the case of aggregates in the absence of hydrolysis in which a shear rate of the same order of magnitude would at most induce slow tumbling. Such low \( Wi \) have an appreciable effect at high hydrolysis rates because the flow need only disperse individual beads rather than long polymeric chains.

CRediT authorship contribution statement

Javor K. Novev: Methodology, Software, Formal analysis, Investigation, Data curation, Writing - original draft, Writing - review & editing, Visualization. Amin Doostmohammadi: Conceptualization, Formal analysis, Writing - review & editing, Funding acquisition. Andreas Zöttl: Methodology, Software, Writing - review & editing, Funding acquisition. Julia M. Yeomans: Conceptualization, Formal analysis, Resources, Writing - review & editing, Funding acquisition, Supervision, Project administration.

Conflict of interest

There are no potential conflicts of interest.

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Appendix A. Supplementary data

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References


