Research paper

Characterisation of human saliva as a platform for oral dissolution medium development

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ABSTRACT

Human saliva is a biological fluid of great importance in the field of dissolution testing. However, until now, no consensus has been reached on its key characteristics relevant to dissolution testing. As a result, it is difficult to select or develop an in vitro dissolution medium to best represent human saliva. In this study, the pH, buffer capacity, surface tension, viscosity and flow rate of both unstimulated (US) and stimulated (SS) human saliva were investigated in order to provide a platform of reference for future dissolution studies using simulated salivary fluids. Age and gender related differences in a sample size of 30 participants for each parameter were investigated. Significant differences were established between US and SS for all characteristics except surface tension. Therefore, the requirement for using two simulated salivary fluids should be considered when developing an oral dissolution model.

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1. Introduction

The oral cavity as a dissolution site is often overlooked due to rapid oral transit as conventional dosage forms are swallowed. However conventional oral formulations such as tablets and capsules are of limited application in the paediatric and geriatric population and alternative oral dosage forms which reside in the mouth for a significant time are increasing in popularity [1]. In addition, adult dosage forms which can be taken “on the move”, without the co-administration of water are also gaining interest [2]. Many alternative formulations, such as oral films, sublingual and buccal tablets and orally disintegrating tablets rely on dissolution or disintegration in saliva. On the contrary, taste masked oral dosage forms often aim to reduce drug dissolution in saliva in order to prevent contact between the unpleasant tasting active pharmaceutical ingredient (API) and the taste buds [3]. Saliva therefore plays a critical role in the dissolution and performance of these formulations. However, there is no consensus on the key characteristics of human saliva which may affect dissolution, and as a result, it is difficult to select or develop an in vitro dissolution medium to best represent human saliva in the evaluation of these dosage forms.

A number of parameters can be considered as highly influential on in vitro dissolution. The pH, buffer capacity and surface tension are identified as some of the most important factors [4]. Additionally, viscosity is considered in many cases [5]. Furthermore, Wang et al. described biorelevant dissolution and suggested consideration of pH, buffer capacity, surface tension and viscosity of the medium to be paramount for biorelevant dissolution testing (along with non-medium related hydrodynamic factors such as volume, flow, agitation and apparatus) [6]. The importance of these particular parameters is evident as similar approaches have been adopted in the characterisation of other gastro-intestinal fluids [7–13]. Extensive research has been carried out over the past few decades in the design and development of, and application of, biorelevant media representing other gastro-intestinal fluids in both the fed and fasted states [11,14–20]. However, saliva remains less well characterised. Without knowledge of these crucial medium parameters in human saliva, it would prove impossible to develop or select a biorelevant simulated salivary fluid for dissolution studies. This paper therefore aims to address the gap in the characterisation of human biological fluids through the investigation of key parameters of saliva.

The pH of a dissolution medium is important since it affects ionisation of the API, according to the Henderson–Hasselbalch equation, and ionisation is directly linked with the aqueous solubility...
of an API [21]. Of equal importance therefore is the ability of the medium to resist changes in pH as an acidic or basic drug begins to dissolve, i.e. the medium's buffer capacity. This was demonstrated by Tsume et al. [22] who performed dissolution experiments in media of different buffer capacities with the acidic drug ibuprofen and found that when the buffer capacity was low, the pH decreased to a greater extent as dissolution proceeded, which hindered the rate and extent of further dissolution.

The pH of human saliva has been described previously, with varying results in the wide range of 5.3–7.8, depending on the stimulation state [10, 23]. In most studies, either unstimulated saliva (US) or stimulated saliva (SS) was investigated, but not both [24, 25]. Additionally, studies in which the pH of both types of saliva was investigated generally had a small number of participants, or focussed on just one or two salivary characteristics relevant to dissolution testing, such as Bardow et al. [26] who investigated only the pH and buffer capacity. We aim to address this issue by characterising the pH of both US and SS, as well as other key parameters within the same sample.

Buffer capacity has been investigated in numerous studies. However, in most cases, the experimental design employed does not allow one to draw conclusions about the actual buffer capacity value or range. Literature values are reported in different ways. Some research groups have simply quoted the buffer capacity to be high, medium or low, without providing any actual value [27]. Thus one cannot draw direct comparisons between studies. Some researchers simply state the bicarbonate concentration of saliva samples to infer buffer capacity [28]. Furthermore, in some cases, buffer capacity is quoted in mmol L\(^{-1}\) pH\(^{-1}\) [26, 29]. The lack of similarity in experimental design has led to inconclusive findings regarding the buffer capacity of saliva. This research aims to address these issues by assessing the buffer capacity of saliva using similar experimental design to that used for other gastro-intestinal fluids [12, 13, 16, 20] to allow for comparison.

Viscosity is another key parameter affecting dissolution. A high viscosity medium would increase the thickness of the boundary layers (h) and decrease the diffusion coefficient (D) according to the Noyes–Witney dissolution model, thus reducing the drug dissolution rate compared with a medium of lower viscosity [30]. Despite viscosity of stimulated and unstimulated whole human saliva being evaluated by several research groups, no consensus has been reached on human saliva viscosity due to differences in experimental conditions. For example, in a review by Schipper et al. [31] viscosity of unstimulated whole saliva was found to be 1.5–1.6 mPa s over a shear rate of 1–300 s\(^{-1}\) in one study [32]. However another study found it to range from 3.8 to 8.8 mPa s at a single shear rate of 90 s\(^{-1}\) [33] and a viscosity of 100 mPa s was recorded at a shear rate of 0.02 s\(^{-1}\) in another study [34] within this review. Research groups used different shear rates, temperatures and types of rheometer and often small sample sizes. This research aims to address these issues by using physiological temperature and assessing viscosity across a wide range of shear rates.

It is well known that the surface tension of the medium also affects the rate of dissolution [9]. A high interfacial tension reduces wetting of the drug particles and reduces the rate of dissolution. Wetting can be improved by the addition of surfactants, reducing interfacial tension and increasing the rate of dissolution, and it is a common practice to add surfactants to dissolution media [35]. Although many studies have investigated the film forming properties of saliva, as well as salivary pellicle thickness and composition [36], few studies have focussed on the surface tension of whole human saliva [23]. Literature regarding the surface tension of saliva uses variable experimental designs including different temperatures and sites in the oral cavity, and often small or non-specified numbers of participants [37, 38]. Further clarification of this parameter is therefore required, using a sufficient number of samples and physiologically relevant temperature.

Despite not directly affecting media choice and composition, salivary flow rate is an important factor when developing a biorelevant dissolution model [6]. The volume available for dissolution, or flow rate, should reflect physiological conditions since this affects the concentration gradient of solvated API molecules and saturation of the bulk fluid. Salivary flow rate has been investigated; however, most groups investigated either US [24] or SS [39]. Since inter-individual variation is so vast in these studies, flow rate should be considered for US and SS in the same individual to allow accurate comparison of stimulation states.

Many research groups performing dissolution tests modelling the oral environment have not taken into account the key characteristics described above and have used simple media such as water or phosphate buffered saline (PBS) [3, 40]. The most likely reason for that is the lack of availability of sufficient data on the characteristics of human saliva. In addition, many commercial [41] and literature formulations [42] for simulated salivary fluids (SSF) exist. In fact, one article in the year 2000 identified 60 artificial salivas of differing compositions. Whilst it is possible that some of these 60 compositions may be appropriate for dissolution studies, information regarding parameters key to dissolution testing is not available [43]. It is also difficult to select the single most appropriate SSF without a clear understanding of human salivary characteristics, and therefore these SSFs are rarely used in dissolution testing. Saliva is a complex mixture containing 99% water, but additionally contains numerous electrolytes, small organic molecules such as hormones and glucose, and proteins such as immunoglobulins, enzymes and glycoproteins, which may have a huge impact on dissolution [44].

Therefore, the aim of this work was to characterise stimulated and unstimulated human saliva for the key characteristics relevant to dissolution to provide a platform of reference for the future selection or development of oral dissolution media that would be representative of human saliva. The saliva flow rate was assessed in this work to aid development of oral dissolution models. Age and gender related differences were also investigated for each parameter. To the best of our knowledge, this is a first work in which the key parameters relevant to drug dissolution – pH, buffer capacity, viscosity, surface tension and flow rate – are assessed simultaneously for both stimulated and unstimulated whole human saliva with a sufficient number of participants to draw statistically meaningful conclusions.

2. Methods and instrumentation

2.1. Human volunteers

All saliva samples were collected in accordance with ethical approval number R12122013 SoP TTTFS, Faculty of Medicine and Health Sciences Research Ethics Committee, Queens Medical Centre, Nottingham University Hospitals. Participation was voluntary and informed written consent was obtained. All data were held in accordance with the Data Protection Act.

Participants were recruited from the University of Nottingham and were healthy adult volunteers. Exclusion criteria included chronic or acute illness in the past 3 months, cold or flu symptoms, oral health concerns and any medication, with the exception of contraception. Participants were asked not to eat, drink, smoke or use oral hygiene for 2 h prior to donation. Donations took place at approximately 3 pm in the afternoon to avoid diurnal salivary changes. The study group demographics are shown in Table 1. The study group was mostly Caucasian (26 of 30 participants).
2.2. Saliva collection and flow rate

Participants were asked firstly to donate an unstimulated saliva sample by draining their saliva via a sterile disposable funnel into two 15 mL polypropylene sterile graduated centrifuge tubes (Greiner Bio-One, UK) and one 1.5 mL polypropylene graduated microcentrifuge tube (Sarstedt, UK). Samples were collected in 3 different vessels to allow for separate defrosting for each characterisation. For each type of saliva (US and SS), the following was collected: 10 mL for buffer capacity, 6 mL for viscosity and 1.5 mL for surface tension measurement. This allowed for a slight excess for each measurement. The time taken to donate each volume was recorded using a stopwatch and the exact volume was used to calculate flow rate for each sample.

Participants then donated a stimulated saliva sample. For this, they were asked to continually chew a 5 cm × 5 cm piece of Parafilm® [25,26,45,46] and repeat the donations following this stimulation. Stimulation was controlled by regulating the size of the piece of Parafilm®, and the same volume of saliva was collected each time. Samples were immediately tested for pH, flash frozen in liquid nitrogen and then stored at −80 °C until being defrosted for characterisation. No significant difference in pH was observed between fresh and defrosted samples (paired t-test).

2.3. Viscosity

A Modular Compact Cone-Plate Rheometer MCR 302 (Anton Paar GmbH, Germany) was used. The cone used was a CP50-2-SN30270 with diameter 49.972 mm, angle 2.016°, truncation 211 μm. Analysis was carried out at 37 °C. 8 points per decade were used for 3 decades with shear rate increasing logarithmically from 1 to 1000 s⁻¹. A total of 25 points were made, 1 point per minute. Rheoplus analysis software (Anton Paar GmbH, Germany) was used. The sample volume was 1.2 mL. Saliva was analysed in triplicate for each participant for US and SS. A range of shear rates was used to assess whether saliva exhibits non-Newtonian behaviour. A shear rate of 4 s⁻¹ corresponds to movement of particles across the tongue, 60 corresponds to swallowing and 160 s⁻¹ to speech, whilst shear rates of 10–500 s⁻¹ have been proposed to reflect the shear during eating. We therefore used 1–1000 s⁻¹ to encompass values that are likely to be presented in the oral cavity [47].

2.4. Surface tension

A DSA 100 Drop Shape Analyser with DSA 4 software (Kruss GmbH, Germany) using pendant drop method for surface tension analysis with Laplace–Young computational method was employed. Temperature was set to 37 °C using an MB-5 heat circulator (Julabo GmbH, Germany) with water bath. Measurements were taken immediately after droplet formation. Samples were measured with 5 replicates.

2.5. pH

An S220 seven compact pH/ion meter was used with InLab Science Pro electrode (SI 343 071, Mettler Toledo, Switzerland). The pH meter was accurate to ±0.002 pH units and a 3 point calibration was used at pH 4, 7 and 10. The pH of human saliva was measured immediately after collection prior to freezing the samples in liquid nitrogen and storing at −80 °C. The pH was measured in triplicate for each participant for both US and SS.

2.6. Buffer capacity

A 4 mL saliva sample was allowed to warm to 37 °C in the test tube. Temperature was maintained using a water bath in which the test tube for titration was placed. The beaker was placed on an RCT basic hotplate stirrer (JKa Works GmbH, Germany) with temperature probe. Initial pH was tested. The sample was then titrated with 0.01 M HCl at 37 °C until a decrease in pH of 1 unit was observed. Buffer capacity in mmol H⁺/L saliva was calculated from the volume of acid added. Stirring speed was set such that the added HCl was adequately mixed throughout the bulk of the sample without forming a vortex. A 100 mm × 23 mm B19/26 glass test tube was used (supplied by Scientific Glassware Supplies, UK). Human saliva was analysed in duplicate for each participant for each type of saliva.

2.7. Statistical analysis

Prior to experiments, the number of study participants was determined using a power calculation [48]. With 80% power and a level of significance of p < 0.05, 95% significance, a sample size of 8–34 participants is sufficient to detect small to very large differences, with sample size being inversely proportional to the difference to be detected.

Following the completion of the experiments, the normal distribution of the results in each group was tested using a D’Agostino & Pearson omnibus normality test. Where two normally distributed groups were compared, a t-test was used (either paired or unpaired). If one or both groups were not normally distributed, a Wilcoxon matched pairs test was used for paired samples, and a Mann–Whitney U-test was used for unpaired samples. p < 0.05 was considered significant in all cases.

3. Results

3.1. Viscosity

The viscosity of US and SS are described in Fig. 1. SS was shown to have a lower viscosity, and a statistically significant difference in viscosity was observed between US and SS at every shear rate recorded with p < 0.0001 (Wilcoxon matched pairs test).

![Fig. 1. The viscosity (mean ± S.D.) of US and SS (n = 30, triplicates) at different shear rates. A statistically significant difference in viscosity was observed between US and SS at every shear rate recorded (p < 0.0001, Wilcoxon matched pairs test). (For interpretation of the references to colour in this figure legend, the reader is referred to the web version of this article.)](image-url)
A statistically significant difference (Mann–Whitney U-test, \( p < 0.05 \)) in US viscosity was observed between males and females at 5 shear rates in the lower shear rate range, with male saliva showing higher viscosity (Fig. 2A). This trend appears to continue across the remainder of the viscosity profile. Practically no difference was observed for SS between male and female groups, with a statistically significant difference (Mann–Whitney U-test, \( p < 0.01 \)) observed at just one shear rate (Fig. 2B).

The viscosity of US was significantly higher for the age group 28–35 compared to 20–27 at 3 shear rates (Mann–Whitney U-test, \( p < 0.05 \), Fig. 3A). This trend also appears to continue across the rest of the viscosity profile. For SS, no significant difference was observed between the two age groups (Fig. 3B).

### 3.2. Surface tension

The surface tension for US and SS is shown in Fig. 4. The surface tension of US was very similar to SS, with no significant difference observed between the two types of saliva (paired t-test). Note the variability between individuals in surface tension of saliva was very low.

The surface tension of human saliva (Table 2) showed no significant difference between males and females for US or SS. In addition, no significant difference in surface tension of human saliva was observed between different age groups for US or SS.

### 3.3. pH

SS had a higher pH than US (Fig. 5) and a statistically significant difference was observed between the two groups according to a paired t-test (\( p < 0.0001 \)).

### 3.4. Buffer capacity

The buffer capacity was found to be significantly different for US and SS (paired t-test, \( p < 0.0001 \)), with SS having a much greater buffer capacity, as shown in Fig. 6.
No significant difference in buffer capacity was observed for US between males and females. However, a significant difference in buffer capacity was observed for SS between males and females (unpaired t-test, \( p < 0.05 \)). No significant difference in buffer capacity was observed between different age groups for US or SS (Table 4).

3.5. Flow rate

As anticipated, the flow rate of SS was significantly greater than US, shown in Fig. 7 (paired t-test, \( p < 0.0001 \)). No significant difference in flow rate was observed between males and females for US or SS. Similarly, no significant difference in flow rate was observed between age groups for US or SS (Table 5).

### 4. Discussion

4.1. Viscosity

Human saliva was found to be non-Newtonian across the range of shear rates applied. The shear rates tested are likely to be in the range observed in the oral cavity since it has been suggested that a
shear rate of 4 s⁻¹ corresponds to the movement of particles across the tongue whilst 60 s⁻¹ and 160 s⁻¹ correspond to swallowing and speech respectively [34,47]. Furthermore, shear rates between 10 and 500 s⁻¹ have been proposed to mimic the range of shear rates in the mouth during eating [49]. US was shown to have a higher variability, with a greater relative standard deviation observed for US than SS at all shear rates measured, with the exception of the very lowest shear rate.

SS’s lower viscosity is proportional to its higher flow rate, leading to an increased aqueous content, and a lower concentration of mucins: glycoproteins with a polypeptide backbone and oligosaccharide side chains which are thought to be responsible for the viscosity of saliva [50]. It has been suggested that this is due to SS originating predominantly from different salivary glands compared to US [33]. SS has been suggested to have a larger proportion of parotid secretions. However, mucins are mainly secreted from the sublingual, submandibular and palatal glands [33]. Indeed, it is well documented that secretions from the main salivary glands have differing mucin proportions and thus differing viscosities. In some cases, parotid saliva has actually been shown to demonstrate Newtonian behaviour, further reinforcing the link between mucin presence and shear thinning behaviour [31,51].

In human saliva, there are two main types of mucin present: a high molecular weight (MW) mucin, MUC5B (MW 2–40 MDa), and a low molecular weight mucin, MUC7 (MW approx. 150 kDa). The molecular structure of mucin is discussed in detail elsewhere by Haward et al. [20] One study investigated which of these types of mucin is responsible for modifying the viscosity of saliva. They established that MUC5B concentration increased linearly with viscosity, but MUC7 did not, thus it is likely that MUC5B is responsible for the viscosity of saliva [22].

The results obtained in this study correspond well with other reports regarding the viscosity of US and SS since other research groups found SS to be of lower viscosity [46,51]. The actual viscosity values for US and SS in the literature vary depending on the type of viscometer, shear rates and temperature used. However, similar to other reports [50–52], we also observed non-Newtonian behaviour for human saliva. This is thought to be attributed to the destruction of the mucin network within the samples which undergo an irreversible breakdown upon shearing [31].

4.3. pH

The higher pH in the SS group can be attributed to differences in electrolyte composition, including a greater bicarbonate concentration in SS [62]. The pH of saliva is modified as it travels through the duct system within salivary glands by the secretion and reabsorption of electrolytes, depicted in Fig. 8. Initially, an isotonic fluid solution media pH values to reflect these findings. As fluid travels along the duct, reabsorption of some ions such as sodium and chloride, and secretion of others such as bicarbonate and potassium occur, until a hypotonic solution is released from the duct [44].

Each acinus may contain only serous cells, mucous cells or both. Serous secretions are rich in electrolytes and enzymes, whereas mucous secretions are rich in glycoproteins. The parotid gland has predominantly serous secretion. Upon stimulation of saliva, there is a greater parotid gland output, thus a greater release of bicarbonate rich serous secretion. This coincides with a lower mucin concentration for SS as discussed previously [44].

Literature stating the pH of human saliva reports variable values that range from 5.3 to 7.8 depending on the stimulation state [23–25]. We found US and SS to be within this range, with mean values of 6.97 and 7.40 for US and SS respectively (range US: 6.49–7.28, range SS: 6.96–7.69). It would be advisable to tailor dissolution media pH values to reflect these findings.
4.4. Buffer capacity

The greater buffer capacity of SS can be attributed to the higher bicarbonate concentration. Bicarbonate contributes approximately 80% of the buffering capacity of human saliva [63], and is found in higher concentrations in SS due to the higher proportion of parotid gland secretions [27]. It should be noted that unlike pH which was measured immediately upon collection, buffer capacity was measured after flash freezing and short term storage at −80 °C. The bicarbonate buffer is a dynamic system and in liquid saliva samples, carbon dioxide may be lost from the system. Although we do not anticipate the buffer capacity to alter as a result of freezing, this could be considered a limitation of the study.

A direct comparison with other literature is challenging due to methodological differences. Nevertheless, the approach used here was also used by Bardow et al. [26] who found the buffer capacity to range from 3.1 to 6.0 mmol H⁺/L of saliva in US and 3.3 to 8.5 mmol H⁺/L of saliva in SS depending on the pH. This is comparable to our values, since we found mean values to be 5.93 and 8.41 mmol H⁺/L of saliva for US and SS respectively. In both cases, SS buffer capacity is higher than US. Despite methodological differences, this was also true for other literature [27]. However, we found buffer capacity to be highly variable for both US and SS, with relative standard deviation being 30.29% and 24.08% for US and SS respectively. This demonstrates a high inter-individual variation, which should be taken into account when designing a dissolution medium.

4.5. Flow rate

The increased flow rate for SS results from the parasympathetic response to Parafilm® chewing which increases saliva output from the salivary glands, in particular the parotid gland. Inter-individual variability was high, with relative standard deviation being 41.0% and 47.5% for US and SS respectively. In this study, US flow rate ranged from 0.23 to 1.10 mL/min with a mean value of 0.58 mL/min whilst SS flow rate ranged from 0.43 to 3.45 mL/min with a mean value of 1.51 mL/min.

Literature is also highly variable, with one study finding a maximum US flow rate of 2.87 mL/min [64], whilst mean SS flow rate was quoted to be just 0.9 mL/min in another study [65]. Across literature, salivary flow rate has been quoted to range from 0.05 to 7.0 mL/min [24,66].

It is known that saliva undergoes diurnal changes in flow rate [66], and since a higher flow rate was associated with a higher pH, higher buffer capacity and lower viscosity in our study, the time at which saliva is collected may affect many of the salivary parameters investigated. Thus, the time of collection was controlled and 3 pm was chosen for practical reasons.

There are three main mechanisms of salivary stimulation: mechanical, gustatory and olfactory [44]. Dissolution testers should consider whether the dosage form may stimulate saliva. Crucially, the presence of a dosage form in the oral cavity such as an orally disintegrating tablet or oromucosal formulation may stimulate the release of saliva and therefore it may be prudent to consider both US and SS when modelling the oral cavity. Given that the flow rate and many other parameters are so variable for human saliva, this reinforces the requirement to model both the US and SS states since a single set of test conditions is unlikely to represent the range of salivary scenarios observed.

4.6. Effect of age and gender on salivary parameters

We observed significant gender and age related differences in viscosity of US in the low shear rate region wherein viscosity was found to be higher for males and the older age group (even with the relatively narrow age range of volunteers). This low shear rate region may require further investigation as statistical differences between demographic groups are only seen in this region. Furthermore, when designing biorelevant dissolution media, this low shear rate region should be modelled accurately. Little is understood about the effect of age and gender on saliva viscosity. Humphrey and Williamson [44] claim that mucin concentrations decrease with age, but also state that secretory hypofunction is not a normal age related phenomenon.

No differences were observed in surface tension for any demographic group. Similar to viscosity, little research has been carried out in this area. Kazakov et al. [23] found that equilibrium surface tension decreased with age, whereby age 5–9 > 10–15 > 40–55 years. However, surface tension in the over 55 years group began to increase so a linear relationship with age was not established. The effect of gender was also not considered in that study. Conversely, extensive literature exists detailing the influence of age or gender on flow rate. Despite this, age and gender related effects remain unclear due to conflicting reports [67]. In this study, no significant differences in flow rate were observed between males and females, or between the two age groups. Accordingly, other researchers also found flow rate was not affected by age [68] or gender [26]. However, some literature suggests that female gender correlates with lower flow rate [28,69] which may be attributed to smaller salivary glands and a lower body mass index (BMI) [70,71]. Additionally, increased age has been reported to correspond with lower flow rates in some cases [28,72]. In a review by Whelton [62] decreased salivary flow in older patients is described as being secondary to disease or medication rather than directly due to ageing, and total flow is considered to be independent of age.

No significant differences in pH or buffer capacity were found for any demographic group in this study except for SS buffer capacity, which was found to be higher for males than females. This is in agreement withWikner and Soder [73] who found females had a lower SS buffer capacity. Fenoll-Palomas et al. [28] also found no significant differences in pH, and higher bicarbonate concentration in men than women. However, their findings were based on US only. Conversely, another report states gender had no effect on buffer capacity [26]. pH has been described as higher for males in some studies [69]. Additionally, literature describing the effect of age on pH [69] and buffer capacity [74] reaches no consensus.

In this research, for the first time the effect of both age and gender on salivary key parameters for dissolution testing was investigated. The age and gender related differences observed were not as distinct as the differences between US and SS. Therefore, the development of two different biorelevant dissolution media representing US and SS is strongly recommended, whilst age and gender related differences should be kept in mind and may require further investigation. This is particularly prudent since taste masked and alternative oral formulations are most commonly used in the paediatric and geriatric population. To note, a limitation of this study is the relatively narrow age range employed. This is a result of recruiting unpaid volunteers from within the University. Further investigations of these key parameters in human saliva in a wider age range would be necessary in order to confirm trends seen in the data. However, it is worthy to note that should the paediatric and geriatric population be investigated, it would be inappropriate to make conclusions about each of those populations as a single group compared to the adult population. For example, a single “paediatric dissolution media” or “paediatric model” is unadvisable since a neonate differs greatly in physiology, body mass and pharmacokinetics to an infant or teenager, and salivary parameters may vary greatly too.
5. Conclusions
US and SS were found to be significantly different to each other for pH, buffer capacity and flow rate, with SS being higher for these characteristics. No significant difference was seen between US and SS for surface tension. SS had lower viscosity with significant differences between US and SS observed across all shear rates measured. US and SS were both found to be non-Newtonian. Significant age and gender related differences were observed in some parameters but were not as distinct as differences between US and SS and may require further investigation.

These findings can be used as a platform of reference for the development or selection of future dissolution media representing human saliva. Since SS was found to be significantly different to US and SS and may require further investigation.

Conflict of interest
The authors declare no conflict of interest.

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The authors declare no conflict of interest.