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# 3D printing of edible hydrogels containing thiamine and their comparison to cast gels

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## **Abstract**

In this study, 3% w/v kappa-carrageenan ( $\kappa$ C) and 2% w/v agar were assessed for their suitability for hot extrusion 3D printing (3DP) and compared to cast gels of equivalent composition. Moreover, incorporation of a model active (thiamine) at varying concentrations, was studied for both 3DP and cast microstructures. Rheology and differential scanning calorimetry showed that thiamine (via electrostatic complexation) reinforced the kappa-carrageenan gel network (up to a certain threshold concentration), whereas the agar gel was structurally unaltered by the active's presence. While the  $\kappa$ C-thiamine formulations were printable (within a relatively narrow formulation/processing window), the agar-thiamine systems were not printable via the current set up. Texture profile analysis (TPA) showed that 3DP  $\kappa$ C-thiamine cylinders had a hardness value of  $860 \text{ g} \pm 11\%$  compared to  $1650 \text{ g} \pm 6\%$  for cast cylinders. When compressed they delaminated due to failure between consecutive layers of material deposited during the printing process; light microscopy revealed distinct layering across the printed gel structure. Release tests at  $20^\circ\text{C}$  showed printed gels expelled  $64\% \pm 2.2\%$  of the total active compared to  $59\% \pm 0.8\%$  from the cast gels over six hours. At  $37^\circ\text{C}$  these values increased to  $78\% \pm 2.6\%$  and  $66\% \pm 3.5\%$  respectively. This difference was believed to be due to the significant swelling exhibited by the printed systems. A simple empirical model, applied to the release data, revealed that thiamine discharge from 3DP gels was solely driven by diffusion while ejection of the active from cast systems had both diffusional and relaxation contributions.

## **1. Introduction**

Additive manufacturing, also known as 3D-printing (3DP), is a layer-by-layer production method that uses digital files to create parts and products. While many different areas of industry have adopted it, its uptake as a means of manufacture at the point of production has still not caught up in areas like homes, health care facilities and pharmacies. However, despite this it is still a growth industry worth billions of dollars per year (McCue, 2012). Most research and utilisation has been focused on plastic polymers (Rahim, Abdullah, & Md Akil, 2019), metal (Buchanan & Gardner, 2019), and ceramics (Chen, et al., 2019) with the emphasis being on the small scale production of highly customised items. Other areas of interest have included pharmaceuticals (A. Goyanes, et al., 2017), biotechnology (D. Singh, Singh, & Han, 2016) and prosthesis development (Koprnický, Najman, & Šafka, 2017). The plug and play nature of 3D printing is appealing because laymen are able to connect a printer, load in the printing material and then download one of many designs from the internet, modify it if they choose to and then print it. This allows customisation at the point of demand and modification without the need for additional tooling or moulding.

From the start of the last decade, food 3DP has started to rapidly develop as an area of research. This is due to various factors such as design of complex geometries without

moulds, production of softer foods that mimic the appearance of normal foods for people with conditions such as dysphagia, while reducing production time and skill level of the person producing it as well as increasing repeatability (Kouzani, et al., 2017), and using 3D printing to precisely control ingredient placement and distribution (Diaz, Van Bommel, Noort, Henket, & Briër, 2018). However, development and subsequent uptake of food 3DP in homes and by industry has been inhibited by various issues. The major problem is that 3D printing is still too slow, with larger objects taking upwards of an hour to produce (Lin, 2015). Moreover, food systems are multifaceted, often consisting of several materials in varying ratios, sensory characteristics related to internal microstructure and thermal transition temperature as well as most food materials not being readily extrudable. This is why a lot of research into food 3DP has focused on natively extrudable food such as chocolate (Lanaro, Desselle, & Woodruff, 2019), cheese (Le Tohic, et al., 2018) and dough (Fan Yang, Zhang, Prakash, & Liu, 2018). Some non-natively extrudable food materials tested for 3D printing include fish surimi gel (Wang, Zhang, Bhandari, & Yang, 2018) and fruit (C. Severini, Derossi, Ricci, Caporizzi, & Fiore, 2018). However, most research has been directed into finding a wider range of materials that can be utilised in food 3DP as well as trying to establish an understanding of how their various properties affect their printability.

Hydrocolloid gels (hereafter referred to as hydrogels) are an area of interest in food 3DP with a large body of work directed into investigations regarding their suitability and utility in this field (Kim, et al., 2018; Z. Liu, Zhang, & Yang, 2018; Rutz, Hyland, Jakus, Burghardt, & Shah, 2015). They are considered ideal owing to the fact that many of them are renewable, widely used already in foods and pharmaceuticals and are known to be biocompatible. Hydrogel printing normally involves the cold extrusion of systems that have already set, requiring little, if any, temperature control. Cold extrusion approaches to date have included mixed hydrogels (Kim, Bae, & Park, 2017), single hydrogels (Gholamipour-Shirazi, Norton, & Mills, 2019) and the use of hydrogels as an adjunctive material (Kim, et al., 2019). The second method is the hot extrusion of hydrogels in the sol state, with the sol-gel transition occurring on the printing bed rapidly after being deposited. This has the advantage of being able to print highly viscous gels far more readily, as they would still be in the sol state unlike the gelled samples; this has been highlighted for cold extrusion (Azam, Zhang, Bhandari, & Yang, 2018) which essentially is printing yield stress materials. There exist fewer examples of hot extrusion hydrogel printing such as kappa-carrageenan ( $\kappa$ C) with gelatin (Warner, Norton, & Mills, 2019),  $\kappa$ C by itself (Díaz, et al., 2019) and agar with gelatin (Serizawa, et al., 2014). Another type of 3DP is freeform reversible embedding of suspended hydrogels (FRESH) which involves directly printing the hydrogel into a support bath of a second hydrogel (Hinton, et al., 2015). This allows for a range of intricate structures to be produced that more accurately mimic a cast gel. This process can also be modified to include different support bath materials such as gellan fluid gels (Compaan, Song, & Huang, 2019). This process so often focused on alginate gels loaded with cells being printed, but there is no reason why cold-set hydrogels could not be printed in this manner, with gelatin having been utilised as a support material (Jin, Compaan, Bhattacharjee, & Huang, 2016). However, this process requires the presence of the fluid gel bath, with fluid gels normally requiring equipment capable of delivering shear while controlling temperature such as a rheometer or a pin stirrer vessel (David A. Garrec & Norton, 2012). This makes it more impractical if 3D printers were to become ubiquitous within homes and healthcare settings. Furthermore having laymen handle more complicated tasks such as removing the remaining support fluid gel could also be a barrier to uptake of this method. Whereas, hot extrusion 3DP will produce finalised products that are immediately ready for use and will not require a constant supply of fluid gel.

Owing to the fact that hot extrusion hydrogel printing requires rapid thermal gelation and a high storage modulus (Díaz, et al., 2019).  $\kappa$ C and agar are considered suitable for hot hydrogel 3DP and have been the focus of most studies in this field.  $\kappa$ C is a sulphated polysaccharide that is extracted from red seaweed. When added to water it has the ability to form thermo-reversible gels in the presence of complementary gelling cations (Hermansson, Eriksson, & Jordansson, 1991). Its gelation is believed to occur through the ordering of randomised coils into double helices and then the aggregation of these helices into a polymeric network (Norton, Morris, & Rees, 1984). Agar is a polysaccharide that is extracted from agarophyte seaweeds. Although like  $\kappa$ C it is as able to form thermo-reversible hydrogels, agar does not require any crosslinkers, it forms physical gels simply through hydrogen bridges; and as such it is uncharged. Agar contains two fractions, agarose and agaropectin with only agarose responsible for gelation (Armisen & Gaiatas, 2009). Both of these polysaccharides have many uses in foods as rheology modifiers and gelling agents (Saha & Bhattacharya, 2010). Vitamin B1 also known as thiamine is a water soluble essential vitamin. It is on the WHO list of essential medications and deficiencies can lead to Wernicke-Korsakoff syndrome and may arise from alcoholism or malabsorption (Kril, 1996).

While agar and  $\kappa$ C are widespread within the food sector, they are also used as drug delivery systems as well, in part due to the biocompatibility of these hydrogels (Nayak & Gupta, 2015; Weiner, 1991). Research has gone into modified release oral dosing (Ito & Sugihara, 1996; Picker, 1999), parenteral preparations (Santoro, et al., 2011; Zhang, Tsai, Monie, Hung, & Wu, 2010) and patches applied directly to the skin (Dalafu, Chua, & Chakraborty, 2010). One of the major issues with the centralised, large scale production of medications is that doses are decided on how many people from clinical trials saw the most benefit at that dose. This can lead to people receiving too much or too little of a medicine despite being given a clinically appropriate dose (Cohen, 1999). This is why 3DP of hydrogels is considered suitable to produce customisable delivery vehicles for medicines tailored to the individual patient at the point of delivery (Fina, et al., 2018; Long, et al., 2019). However, 3DP of hydrogels is suited best for smaller batches that require high levels of customisation, whether this is for printing medicines or implant devices (Ventola, 2014). Furthermore there is still an issue with a limited range of materials currently available (Ngo, Kashani, Imbalzano, Nguyen, & Hui, 2018). While studies such as this one aim to address this issue, there is still some way to go. Finally, there will have to be widespread training in order to familiarise medical professionals at the point of production or delivery in the use of 3D printers (Choonara, du Toit, Kumar, Kondiah, & Pillay, 2016; Ventola, 2014).

This study aimed to evaluate the suitability of hydrogels for hot extrusion 3DP and compare and contrast the physical properties of printed gels to cast gels, before assessing them both as release vehicles. Because, there exists little literature on the interactions of thiamine with agar and  $\kappa$ C, and none looking into how thiamine might affect the microstructure of the hydrogels, the first step was to characterise the thermal characteristics of the  $\kappa$ C-thiamine and agar-thiamine hydrogels. Rheology and differential scanning calorimetry were used to determine which formulations were suitable candidates for hot extrusion 3DP.  $\kappa$ C and agar were chosen for printing because of their desirable gelling characteristics. Thiamine was chosen as a model molecule because it has been used in release studies before (Kevadiya, et al., 2010). After suitable thiamine containing hydrogel systems were established then 3D printing under several parameters took place. The printed gels' physical properties was ascertained through texture profile analysis and light microscopy and then compared to cast gels. This highlighted the variances in the structures fabricated between the two production methods. Finally the printed and cast gels underwent release tests to assess their performances as release vehicles in water. This allowed this study to examine physical

differences between printed and cast hydrogels and show compare the performance of 3DP hydrogels to cast gels as drug delivery vehicles.

## **2. Materials and methods**

### **2.1. Materials**

κC, agar and sodium chloride were purchased from Sigma-Aldrich (UK). Thiamine hydrochloride 99% (hereafter referred to as thiamine) was purchased from Alfa Aesar (UK). Sodium hydroxide 1M was purchased from Honeywell and was used for pH adjustment. Milli-Q water was used (Elix® 5 distillation apparatus, Millipore®, USA) for sample preparation. All materials were used without further purifications or modifications.

### **2.2. Hydrogel Preparation**

Samples of κC hydrogels were prepared by dispersing 3% w/v of κC into deionised water. First, a hotplate was set to 80°C and then the water was placed on top of it in a 250mL beaker. A magnetic stirrer was used in order to facilitate dispersion and hydration of the κC into the water. After adding the κC powder into the water, it was left to stir for sixty minutes. Agar hydrogels were produced by placing 2% w/v agar into deionised water. This was then covered and placed in an 800W microwave and heated for ninety seconds until the agar melted into the water. After sixty seconds the microwave was stopped and the beaker was shaken gently. After this the solution was stored on a hotplate set to 70°C and stirred by a magnetic stirrer.

### **2.3. Thiamine hydrogel preparation**

For agar samples containing thiamine, the same methodology used in 2.2 was followed. However, thiamine at the required concentration (0.1%, 1%, 2% and 5% w/v) was added and then the pH was balanced back up to 5.5 using sodium hydroxide 1M and the solution was then held at 70°C until it was ready to be used. This pH was chosen as agar did not undergo acid hydrolysis at this pH (Phillips & Williams, 2000) and the thiamine was still stable at this pH and temperature (Arnold & Dwivedi, 1971). For κC samples containing thiamine, first the thiamine was added to water and then the pH was adjusted to pH 5.5 once more to protect both the thiamine and the κC from degradation. Owing to the fact that sodium ions affect gel strength of κC hydrogels (Hermansson, et al., 1991) the exact amount of sodium hydroxide added for each concentration of thiamine was calculated from the amount of 1M sodium hydroxide added. The amount of sodium ions added to the highest concentration of thiamine was calculated to be 546 mg. Therefore, the 0%, 0.1%, 1% and 2% κC-thiamine gels had sodium chloride added to them to ensure they all contained 546 mg of sodium ions. This ensured any changes to the gel strength alone came from the thiamine and not the sodium ions.

### **2.4. Rheology**

Rheological analysis of the samples were carried out using a modular compact rheometer 302 (Anton Paar, Austria), using parallel 50 mm sandblasted plates or 20 mm serrated plates. A working gap of 1 mm was used for all measurements. All samples were covered with silicone oil around the edges to minimise evaporation during testing. Temperature sweeps were carried out at a fixed frequency of 1 Hz within the linear viscoelastic region. The sweeps ran from 70°C to 20°C except for 5% w/v thiamine and κC which was carried out from 80°C to 20°C owing to the far higher gelling point of the system. This helped to prevent errors whereby the rheometer gave the initial storage modulus ( $G'$ ) as higher than the initial loss modulus ( $G''$ ) despite being above the sol-gel transition temperature. Temperature sweeps were carried out at a scanning rate of 1°C min<sup>-1</sup> to be in line with previous studies (S. Liu & Li, 2016; Tomšič, Prossnigg, & Glatter, 2008). Information from the temperature

sweeps gave data on  $G'$ ,  $G''$  and the phase angle ( $\tan \delta$ ). From this information it was possible to determine the gelling ( $T_{gel}$ ) and melting ( $T_{melt}$ ) temperatures of the gels as the point where  $G'$  and  $G''$  cross over (Djabourov, Leblond, & Papon, 1988).

## 2.5. Micro differential scanning calorimetry

Micro differential scanning calorimetry ( $\mu$ DSC) was carried out using a Seteram MicroDSC 3 evo (Seteram, France). This involved analysis of the thermal transitions of the tested hydrogels. 0.6-0.8 g of sample was loaded into a stainless steel cell. Then the reference cell was filled with the equivalent amount of deionised water  $\pm 0.005$  g. Samples were then subjected to the following measurement conditions used in previous studies (Brenner, Wang, Achayuthakan, Nakajima, & Nishinari, 2013; Iijima, Hatakeyama, & Hatakeyama, 2014).. First, they were cooled to  $0^{\circ}\text{C}$  and held there for sixty minutes. Then they were heated up to  $100^{\circ}\text{C}$  at a scanning rate of  $1^{\circ}\text{C}$  per minute and cooled back down to  $0^{\circ}\text{C}$  at the same rate. Then it was held for another sixty minutes and the cycle was repeated twice more. This gave three heating and cooling curves per run. Each different formulation was tested in triplicate in this manner, giving a total of nine cooling and heating curves per formulation (Brenner, et al., 2013; Iijima, et al., 2014). The temperature range of  $0$ - $100^{\circ}\text{C}$  was chosen because all thermal transitions in the tested systems occur within this range. Transition temperature was taken as the peak of the curve and were obtained by integrating the area below the baseline. Changes in enthalpy ( $\Delta H$ ) were also determined through this method (Iijima, et al., 2014).

## 2.6. 3D printing

The 3D printing system was created from a commercially available printbot simple metal printer which was retrofitted to handle a liquid feed. This involved replacing the components which originally fed the plastic filament into the hot end. The new parts were computer-aided design (CAD) 3D printed parts (4. and 5. in Figure 1) which facilitated the use of a 10 mL syringe (1. In Figure 1) Due to the fact that there was no heating system, the syringe was insulated to prevent temperature loss. This was to maintain the sol state in order to allow the sol-gel transition to occur *in situ* such as with (Warner, et al., 2019). The syringe was then filled with the hot liquid sample and nozzles of several different internal diameters were tested (2. In Figure 1). Printing was carried out at ambient temperature which was set to  $20^{\circ}\text{C}$  by the climate control. Several printing parameters were adjusted depending on the sample such as infill %, print speed, flow %, and layer height. The software used to control the printer was cura software which is freeware available online. Previous works have shown that varying these parameters can have a major impact on the outcome of print fidelity in food systems (C. Severini, et al., 2018; Fanli Yang, Zhang, Bhandari, & Liu, 2018). This proved to be true in this case with many failed prints occurring during parameter optimisation. Printability was assessed through shape fidelity (Chimene, Lennox, Kaunas, & Gaharwar, 2016) and weight uniformity (Goyanes, Buanz, Basit, & Gaisford, 2014). If the printed shapes were close to the computer generated image and were within 5% of the average weight of the printed samples they were considered to be successful. Another risk was premature gelation on the printing bed itself, which led to the nozzle dragging through gelled material owing to the pattern of printing. Conversely, if the sol-gel transition had happened too late then the hydrogel spread across the printing bed and fail to achieve layer by layer build up (Wei, et al., 2015). Layer height has been shown to affect the final print outcome in cold extrusion hydrogel 3D printing (Carla Severini, Derossi, & Azzollini, 2016). This also held true with hot extrusion printing. If the layer height was set too high the hydrogel solution came out in drops, giving broken lines and low quality prints. If it was set too low then the nozzle dragged through the gel, yielding a low quality print. If the bed (3. In Figure 1) was too cold, the first layer set too quickly and then the print might fail as the nozzle might have scraped through the set material. If the bed was too hot, the print was of

low quality as the first layer spread across the build plate due to a failure to set quickly enough. This led to subsequent layers being deposited incorrectly and the final product not properly resembling the CAD shape. Before each print the printer bed level was calibrated manually using a 100  $\mu\text{m}$  gauge. A schematic of the printer is shown in Figure 1.

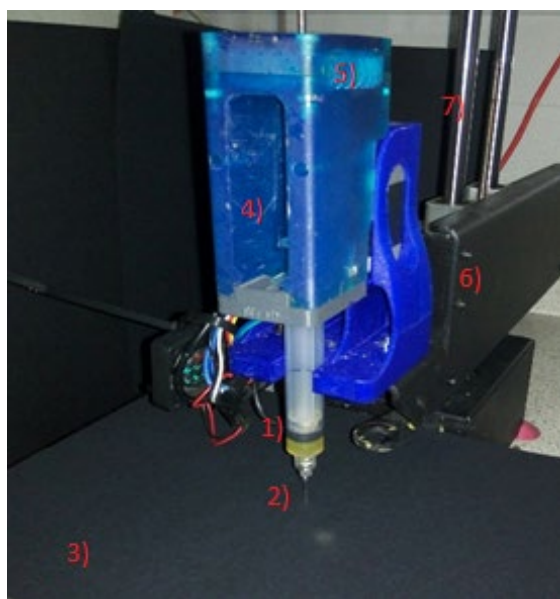


Figure 1: Schematic of the retrofitted printrbot simple metal printer including 1) Syringe to hold liquid feed, 2) Nozzle for extrusion, 3) Temperature controlled printing bed, 4) 3DP bracket to hold syringe, 5) Syringe driver for extrusion, 6) Arm to control movement in the X and Y-axes, 7) Support rods enabling movement in the Z-axis. Printer was connected to and controlled by a computer running cura freeware.

## 2.7. Production of moulds for casting

Moulds were produced by stereolithography 3D printing using a form 2 3D printer (Formlabs, USA). A cube and a cylinder mould were designed by CAD and uploaded to the software, this was then sliced and sent to the printer digitally to print.

## 2.8. Texture Profile analysis

Texture profile analysis (TPA) of the printed samples and the cast control samples was carried out using a TA XT plus Texture Analyser (Stable Micro Systems Ltd. UK) with 30 kg load cell, 3 g trigger force, P/40 (40 mm) cylindrical aluminium probe at a constant speed of 1 mm/s to match previous studies (David A Garrec & Norton, 2013). 12 mm<sup>3</sup> cubes and cylinders of 12 mm height and diameter were printed to be used for testing (see Figure 5.) 12mm<sup>3</sup> cubes and cylinders of 12 mm height and diameter were cast and used as control samples. After printing, samples were tested immediately while, for cast samples, the hydrogel solution was poured into the mould and left to gel at ambient temperature for two minutes; this was approximately equal to the printing time. Cast samples were then immediately evaluated in the texture analyser. Each test was carried out in triplicate. Through compression testing data on hardness and Young's modulus were obtained for the printed and cast samples. Hardness is defined as the peak force during the first compression cycle (Jones, Woolfson, & Brown, 1997). Young's modulus also known as elasticity, is the stiffness of the material calculated through the relationship between stress and strain of the material at low strains (Jones, et al., 1997).

## 2.9. Reflective light microscopy

An optical microscope (DM 2500 LED, Leica®, CH) was used to examine central cross sections of printed and cast hydrogels. The cross sections were obtained by slicing the gels thinly with a scalpel. They were then placed on a glass slide and a cover slip was placed

over the top. The microscope was set to reflective bright field settings and the software included was used to optimise the image. 4 times and 10 times magnification objectives were used. Images were captured using a charge coupled device camera (DFC450 C, Leica®, CH) attached to the microscope.

## 2.10 Release studies

Release studies were carried out using UV-visible spectrophotometry to assess the release of thiamine from the printed gels and compare it to that from cast gels. The gels each contained 2% w/v thiamine. Three cylinders of 12 mm height and 12 mm diameter were printed and each one was placed into a beaker containing 100mL of deionised water. Water was used as a simple, preliminary medium. In the future more complex media will be considered. Cylinders of the same height and diameter were also cast and used as control tests. Owing to thiamine's extremely high water solubility (Pharmacopoeia, 2016) this was an acceptable phase volume to obtain sink conditions (Gibaldi & Feldman, 1967). The beakers of water were put into an Incu-Shake MIDI shaker incubator (Sciquip, UK) at 100 rpm. Release tests were carried out at 20°C in order to test out room temperature for uses other than ingestion and 37°C for *in vivo* testing. Measurements were taken at 0, 5, 10, 15, 30, 60, 90, 120, 180, 240, 300 and 360 minutes and 24 and 48 hours. Determination of the concentration of thiamine within the dissolution medium were carried out using an Orion AquaMate 8000 UV-VIS Spectrophotometer (Thermo Fisher Scientific, UK) set to 235 nm, which was the wavelength at which the thiamine was best detected by the spectrophotometer. 20 µL of dissolution medium was taken with an eppendorf pipette and added to a 1000 µL cuvette. Then 980 µL of deionised water were added to the cuvette and the solution was homogenised in a vortex shaker for 15 seconds to be in line with previous studies (Hansen & Warwick, 1978). This was then placed into the UV-VIS spectrophotometer after it had it calibrated with a blank cuvette containing only deionised water. This gave the concentration of thiamine within the cuvette which was then adjusted to account for the total thiamine released within the dissolution medium. The release profile was calculated from a calibration curve determined by the UV-visible spectrophotometer which had an R<sup>2</sup> value of 0.998. The cuvette was then discarded and 20 µL of deionised water was added into each beaker. This was corrected for when calculating the thiamine concentration following the procedure of (B. Singh, Kaur, & Singh, 1997). All tests were carried out in triplicate.

## 2.11 Modelling of release data

Thiamine release data (up to 60%) were fitted to the model proposed by (Peppas & Sahlin, 1989):

$$\frac{M_t}{M_\infty} = k_1 t^m + k_2 t^{2m} \quad \text{Eq [1]}$$

Where  $M_t/M_\infty$  is the fraction of active released at time  $t$ . The first term ( $k_1 t^m$ ) relates to Fickian effects while the second term ( $k_2 t^{2m}$ ) to relaxational contributions to the release.  $k_1$  is the kinetic constant regarding release from the matrix by Fickian diffusion and  $k_2$  is the kinetic constant for case-II relaxation. Lastly the coefficient  $m$  is the purely Fickian diffusion exponent which is dependent on the shape of the device (Peppas, et al., 1989); the value of the exponent concerning relaxation transport is in theory twice the Fickian exponent ( $2m$ ). The same study further reported that the impact of each of the two mechanisms to the obtained release profile can be assessed by calculating the fractional Fickian (F) and relaxational (R) contributions from:

$$F = \frac{1}{1 + \frac{k_2 t^{2m}}{k_1}} \quad \text{Eq [2]}$$



$$R = \frac{\frac{k_2 t^m}{k_1}}{1 + \frac{k_2 t^m}{k_1}} \quad \text{Eq [3]}$$

### 3. Results and discussion

#### 3.1. Pre-printing thermal characterisation of the hydrogels

Before printing could occur it was important to establish the thermal characteristics of the gel systems. This was crucial as a strong understanding of these parameters is necessary in order to establish whether a material is printable or not. There exists virtually no literature on the effect thiamine has on the thermal characteristics of κC and agar gels. So investigations had to be carried out in order to establish if there were any changes to the gel networks following thiamine incorporation. The sol-gel transition temperature of the thiamine-biopolymer systems was determined using a rotational rheometer. This is in line with previous studies (Hermansson, et al., 1991; Watase & Arakawa, 1968) The results for average  $T_{\text{gel}}$  and  $T_{\text{melt}}$  for 3% κC and 2% agar with 0, 0.1, 1, 2 and 5% w/v thiamine are shown in Figure 2.  $G'$  and  $G''$  for all the examined systems are presented in Figure 3.

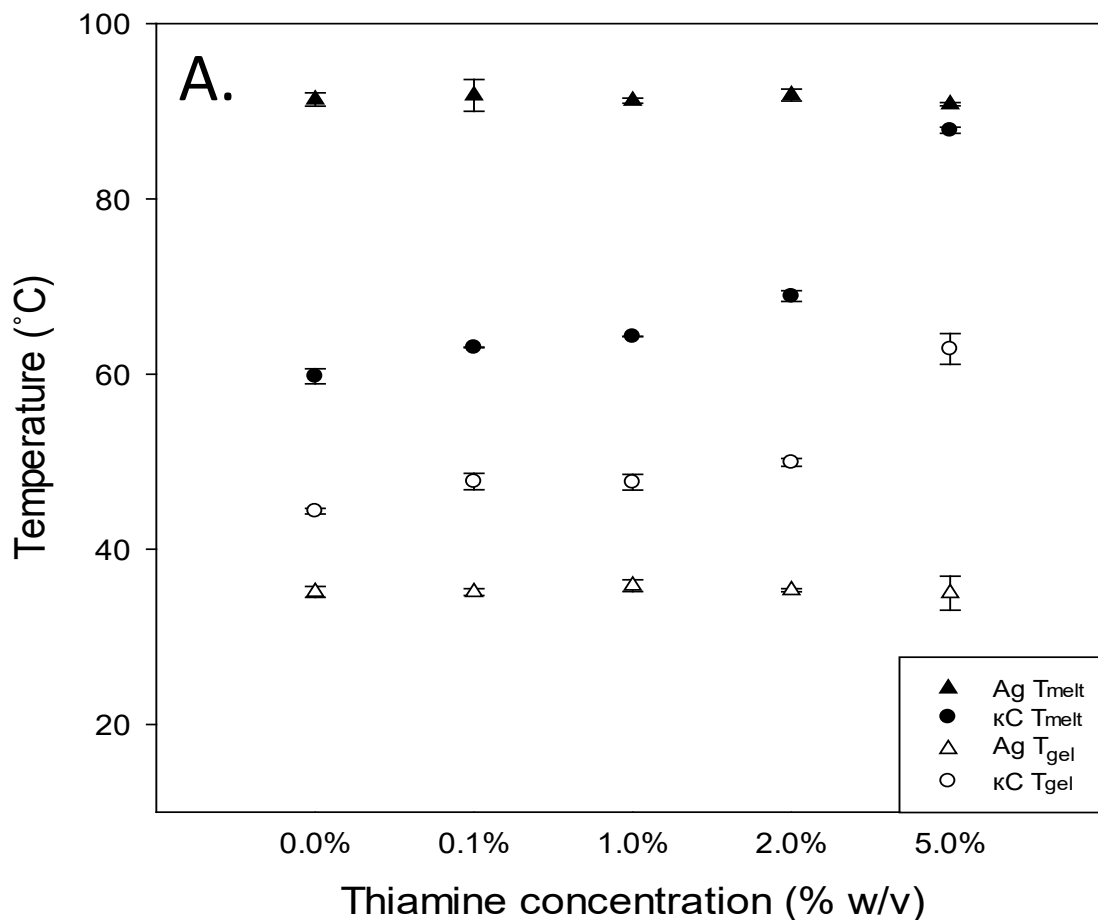


Figure 2:  $T_{\text{gel}}$  and  $T_{\text{melt}}$  of 0, 0.1, 1, 2 and 5% thiamine with 3% κC and 2% agar – Error bars are the standard deviation of the mean.  $n = 3$

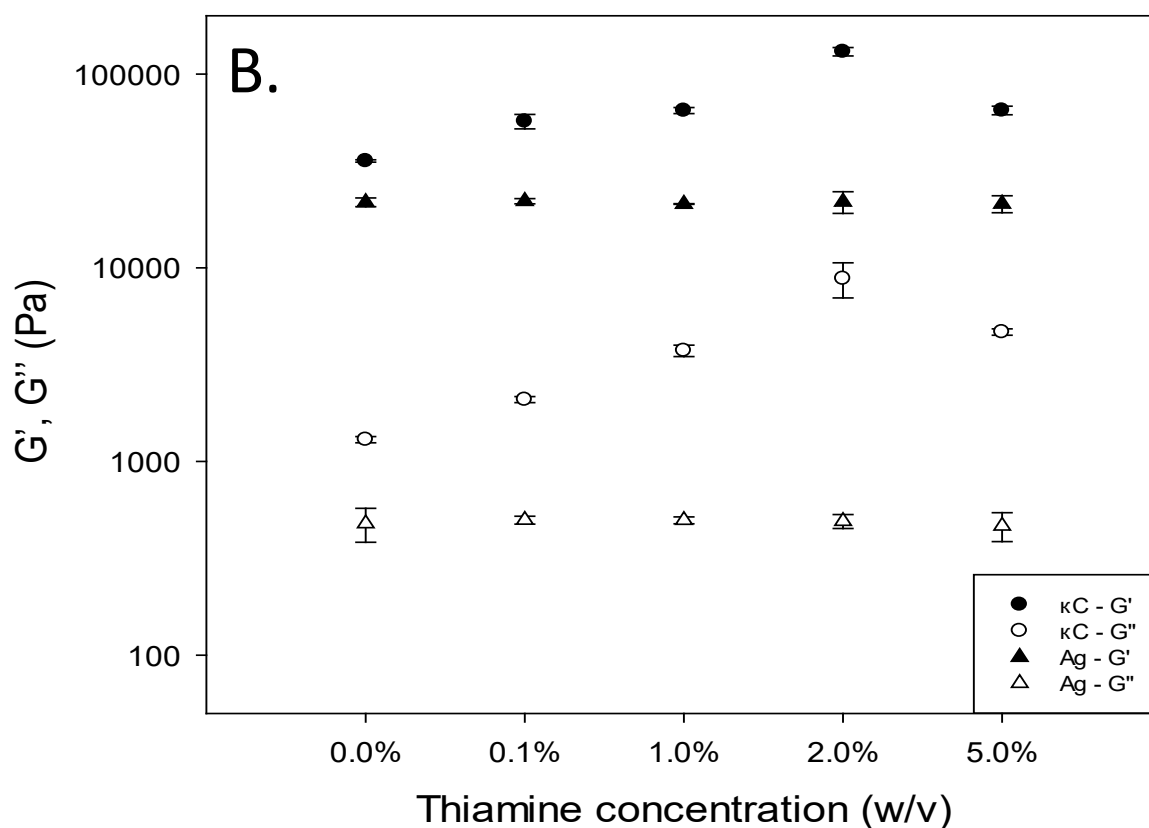
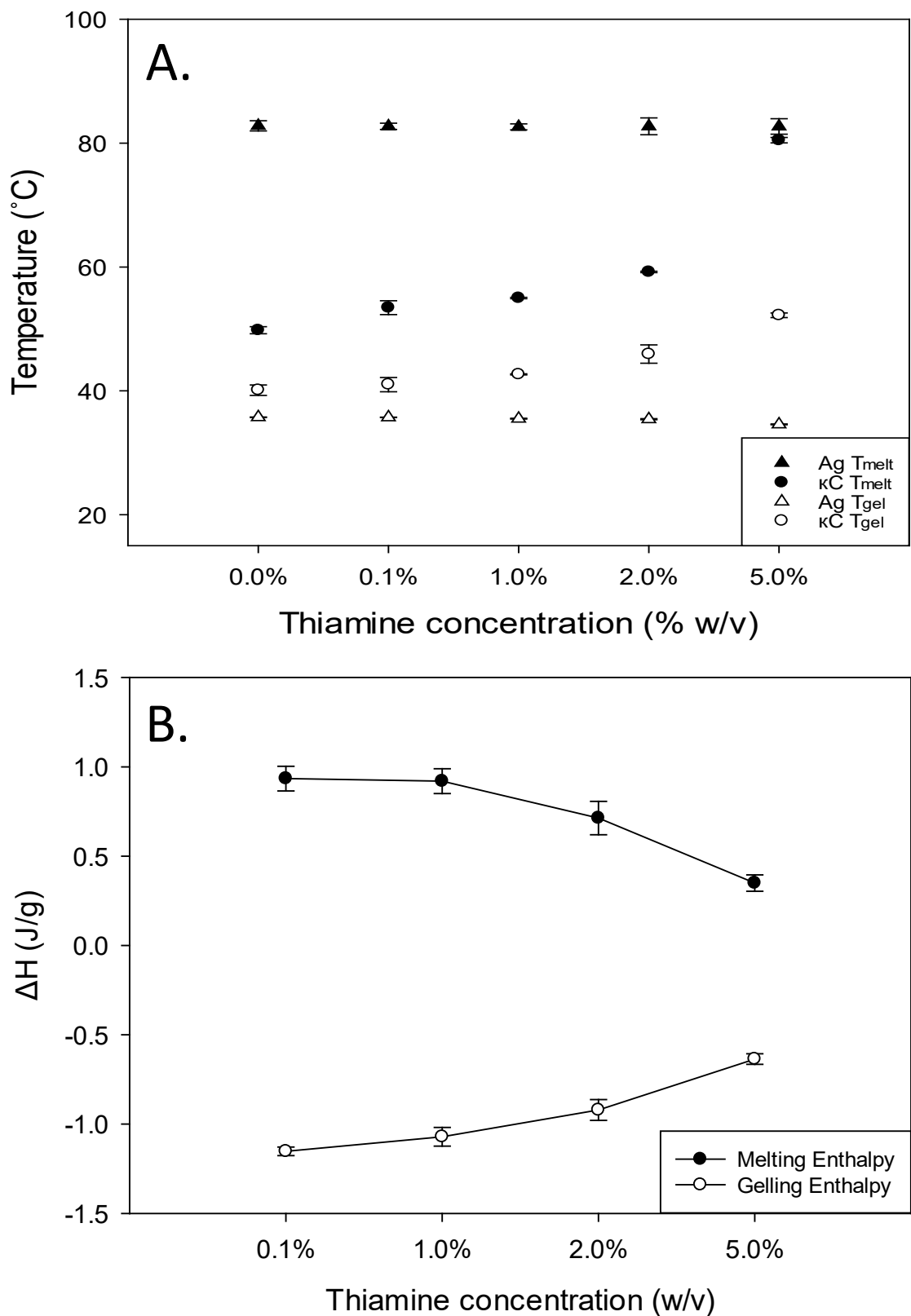


Figure 3:  $G'$  and  $G''$  of 0, 0.1, 1, 2 and 5% thiamine with 3%  $\kappa\text{C}$  and 2% agar

Figures 2 and 3 show that there is no interaction between the agar and the thiamine. As an uncharged molecule, agar gels through hydrogen bonding and doesn't rely on crosslinkers (Tako & Nakamura, 1988). This is reflected in Figures 2 and 3 by the transition temperatures and the moduli of agar remaining constant regardless of thiamine concentration. However, with  $\kappa\text{C}$  and thiamine, as the concentration of thiamine increased there was a linear increase in both  $T_{\text{gel}}$  and  $T_{\text{melt}}$ . This indicates that an interaction between thiamine and  $\kappa\text{C}$  was occurring with the transition temperatures increasing with the concentrations of the active. After dissociating from the hydrochloride salt, thiamine is a cationic molecule and  $\kappa\text{C}$  is an anionic molecule which relies on cationic ions for gelation (Hermansson, et al., 1991). Therefore the results from the  $T_{\text{gel}}$  and  $T_{\text{melt}}$  suggest that the thiamine is complexing with the  $\kappa\text{C}$  and reinforcing the gel network. This phenomenon has been observed with  $\kappa\text{C}$  and other cationic molecules such as surfactants (Grządka, 2015). However, Figure 3 shows that the reinforcement of the gel network through increasing thiamine concentration, peaks at 2% w/v thiamine. At 5% thiamine a decrease in  $G'$  and  $G''$  were observed despite increasing  $T_{\text{gel}}$  and  $T_{\text{melt}}$ . This was probably due to the  $\kappa\text{C}$  becoming saturated with the thiamine, which is a less effective gelling agent than ions such as potassium and sodium. This led to a decrease in the aggregation of double helices which are essential to normal  $\kappa\text{C}$  gelation. The formation of these thiamine- $\kappa\text{C}$  complexes caused charge cancellation and therefore hydrophobic domains which will increase the transition temperatures and inhibit gelation. This has been shown before with cationic compounds reducing gelation of  $\kappa\text{C}$  and even preventing it when they are solely present (Norton, et al., 1984).

However, rheological results alone are not enough to conclusively show that the complexation of  $\kappa$ C and thiamine is occurring. The same systems were tested using a DSC as well. This reaffirmed the transition temperatures and gave information on the gelling and melting enthalpies of the systems as well. Figure 4A shows the  $T_{gel}$  and  $T_{melt}$  for the thiamine-biopolymer systems. Figures 4B and C show the gelling and melting enthalpies.



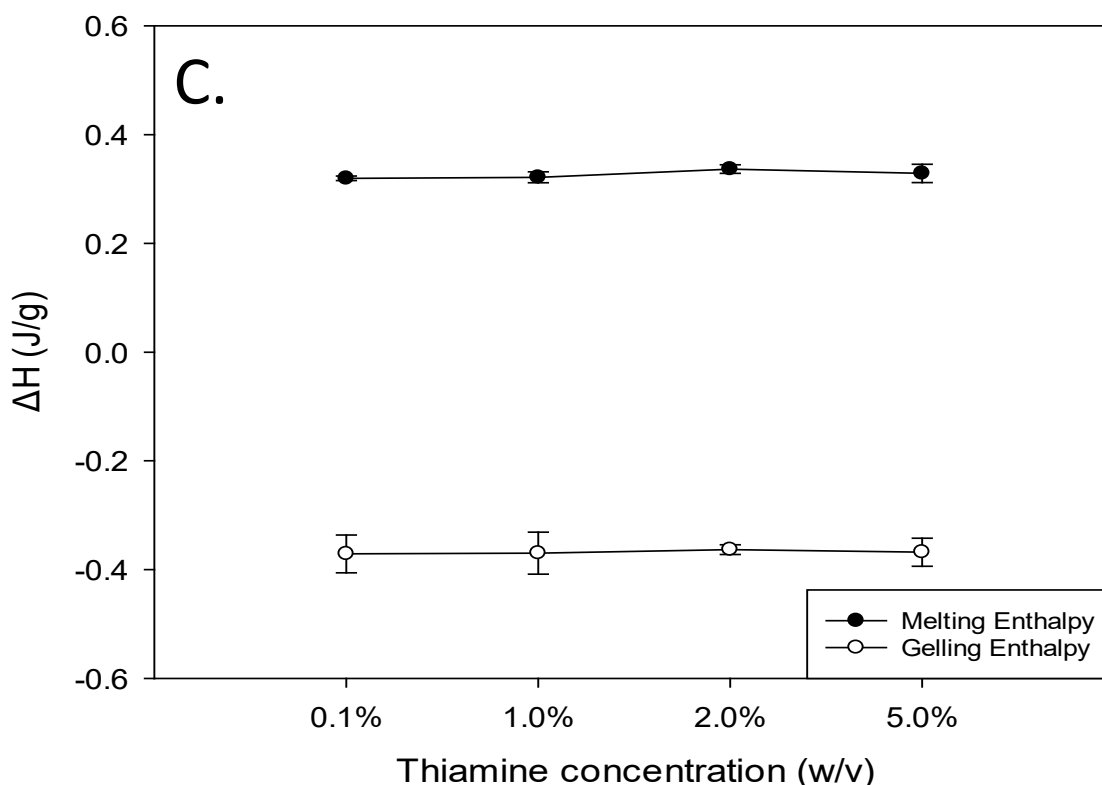


Figure 4: DSC data for the average thermal transition temperature of  $\kappa$ C- and agar-thiamine gels (A) and the gelling and melting enthalpies of  $\kappa$ C-thiamine (B) and agar-thiamine (C) gels.

The  $T_{\text{gel}}$  and  $T_{\text{melt}}$  for the thiamine-biopolymer systems were in agreement with the results from the rotational rheometer. While they do return somewhat different results, this is to be expected owing to the different ways in which they assess the coil to helix transition and vice versa. The rheometer and the DSC are sensitive to different parts of the gelation process of the hydrogels and the results are therefore not obtained from the same segment of the gelling mechanism (Nishinari, 1997). Again, the  $\mu$ DSC results for the thiamine-agar systems confirmed that no interaction is taking place between the agar and the thiamine, with constant transition temperatures and enthalpies recorded regardless of thiamine concentration. However, with the  $\kappa$ C, the thiamine concentration had a negative correlation with the gelling and melting enthalpies. The decrease in enthalpy values indicated that there was a reduction in the amount of free sulphate groups on the backbone available for formation of electrostatic bridges with gelling cations (Rosas-Durazo, et al., 2011). Oversaturation with  $K^+$  ions has been shown to lead to the disruption of  $\kappa$ C cross-linking and prevention of the aggregation of double helices (Thrimawithana, Young, Dunstan, & Alany, 2010), and this phenomenon is believed to occur within these systems. The thiamine- $\kappa$ C hydrogels also became visually more turbid with increasing thiamine concentrations, suggesting the formation of larger complexes which were able to scatter light.

### 3.2. Hydrogel printing

The samples chosen for printing were those that had a higher storage modulus and exhibited rapid solidification (Li, Li, Qi, Jun, & Zuo, 2014). A higher storage modulus has been shown to produce printed products with better shape retention (Costakis, Rueschhoff, Diaz-Cano, Youngblood, & Trice, 2016). The thermal characterisation identified that hydrogels with a 2% thiamine concentration were best suited for 3DP, as it gave the highest storage modulus and gelled rapidly for the  $\kappa$ C. The agar was the same regardless of the

thiamine concentration so 2% was also used. The parameters tested for the 3D printing of  $\kappa$ C-thiamine hydrogels are shown in table 1 below.

*Table 1: A table showing the different parameters tested in the hydrogel 3D printing process*

NS	$H_L$ (mm)	Flow (%)	$T_{PB}$ (°C)	$v_p$ (mm/s)	$T_H$ (°C)	3D Outcome and Comments
18G	1.4	50	40	30	80	Under extrusion – Set too slowly
18G	1.4	60	45	30	75	Under extrusion – Set too quickly
18G	1.4	70	45	20	75	Over extrusion – Set too slowly
20G	1	35	40	20	75	Slight under extrusion – Set properly – nozzle dragged through shape
20G	1	40	40	20	75	Sufficient extrusion – Set properly – nozzle dragged through shape
20G	1.2	35	40	20	75	Slight under extrusion – Set properly
20G	1.2	35	40	20	80	Slight over extrusion – Set properly
20G	1.2	40	40	20	70	Slight under extrusion – Set properly
20G	1.2	40	40	20	75	Sufficient extrusion – Set properly
20G	1.2	40	40	20	80	Over extrusion – Set properly
20G	1.2	45	40	20	75	Slight over extrusion – Set properly
20G	1.2	50	40	20	75	Over extrusion – Set too slowly
20G	1.4	35	40	20	75	Under extrusion – Set properly
20G	1.4	40	40	20	75	Under extrusion – Set properly
20G	1.4	45	40	30	75	Over extrusion – Set properly – nozzle dragged through shape
20G	1.4	50	45	20	80	Over extrusion – Set too slowly
22G	1.4	30	40	20	60	Under extrusion – Set too quickly
22G	1.4	40	45	30	75	Under extrusion – Set too slowly
22G	1.4	50	50	40	80	Over extrusion – Set too slowly

NS: Nozzle size

$H_L$ : Layer height (mm)

Flow: Flow percentage (%)

$T_{PB}$ : Printer bed temperature (°C)

$v_p$ : Print speed (mm/s)

$T_H$ : Hold temperature (°C)

The printing parameters from table 1 that yielded the highest quality and most repeatable prints for thiamine-κC hydrogels were a 20 gauge nozzle, layer height 1.2 mm, flow percentage of 40%, printer bed temperature 40°C, print speed of 20 mm/s and a hold temperature of 75°C.

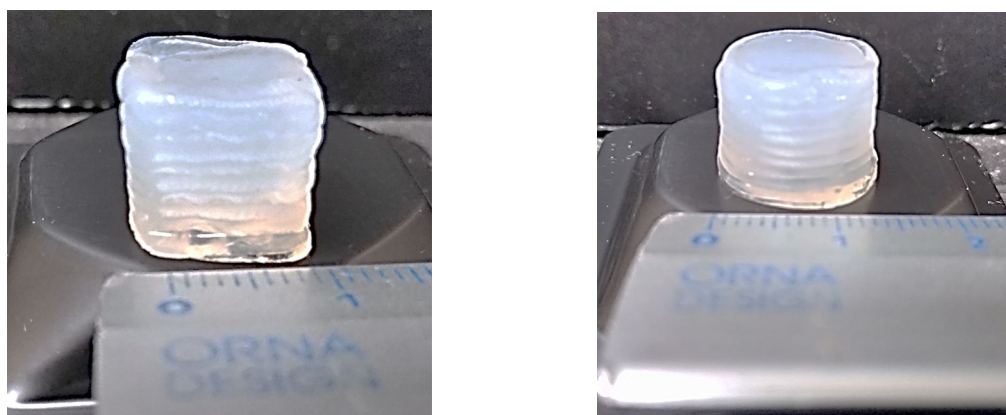


Figure 5: 3D printed 12 mm cube (A) and 12 mm height and diameter cylinder (B) printed using 3% κC and 2% thiamine hydrogel

The agar was unable to successfully print under all tested conditions. This was believed to be because the agar-thiamine hydrogels with their far lower  $G'$  of 20,000 pascals had poorer shape retention compared to the κC-thiamine hydrogels which had a  $G'$  of around 120,000 pascals. Agar's lower  $T_{gel}$  of around 37°C might also have been a contributing factor as well. This meant that there was less of a temperature differential between the  $T_{gel}$  and printing temperature. This led to a slower gelation time and poorer shape fidelity, with the shape unable to hold the weight of subsequent layers. Too much spreading also meant that the nozzle dragged through the hydrogel solution, distorting the shape. The reasons for failure might also have been due to limitations with the printer hardware, as a lack of temperature control on the syringe meant that the hydrogel solution could not be held just above its  $T_{gel}$ . Other modifications such as a cooling fan might also have yielded improved results. Finally changes to the formulation such as increasing the concentration of the agar or addition of an adjunct such as sucrose to increase gel strength and temperature could be successful in future (Normand, 2003). Therefore going forward only thiamine-κC gels were used.

### 3.3. Post-printing texture profile analysis of the hydrogels

TPA of gels is often used to assess their microstructure performance and how this affects specific functionality, including their ability to deliver therapeutic molecules (Özcan, et al., 2009). With the layer by layer nature of 3D printing, the 3D printed structures have a different internal structure compared to a cast/moulded structures (Padzi, Bazin, & Muhamad, 2017). However, there is not much literature that compares 3DP hydrogels to their cast equivalent, and thus TPA was used to begin understanding and characterising some of the internal differences. Figure 4 shows the data obtained for the hardness and Young's modulus of printed and cast cubes and cylinders.

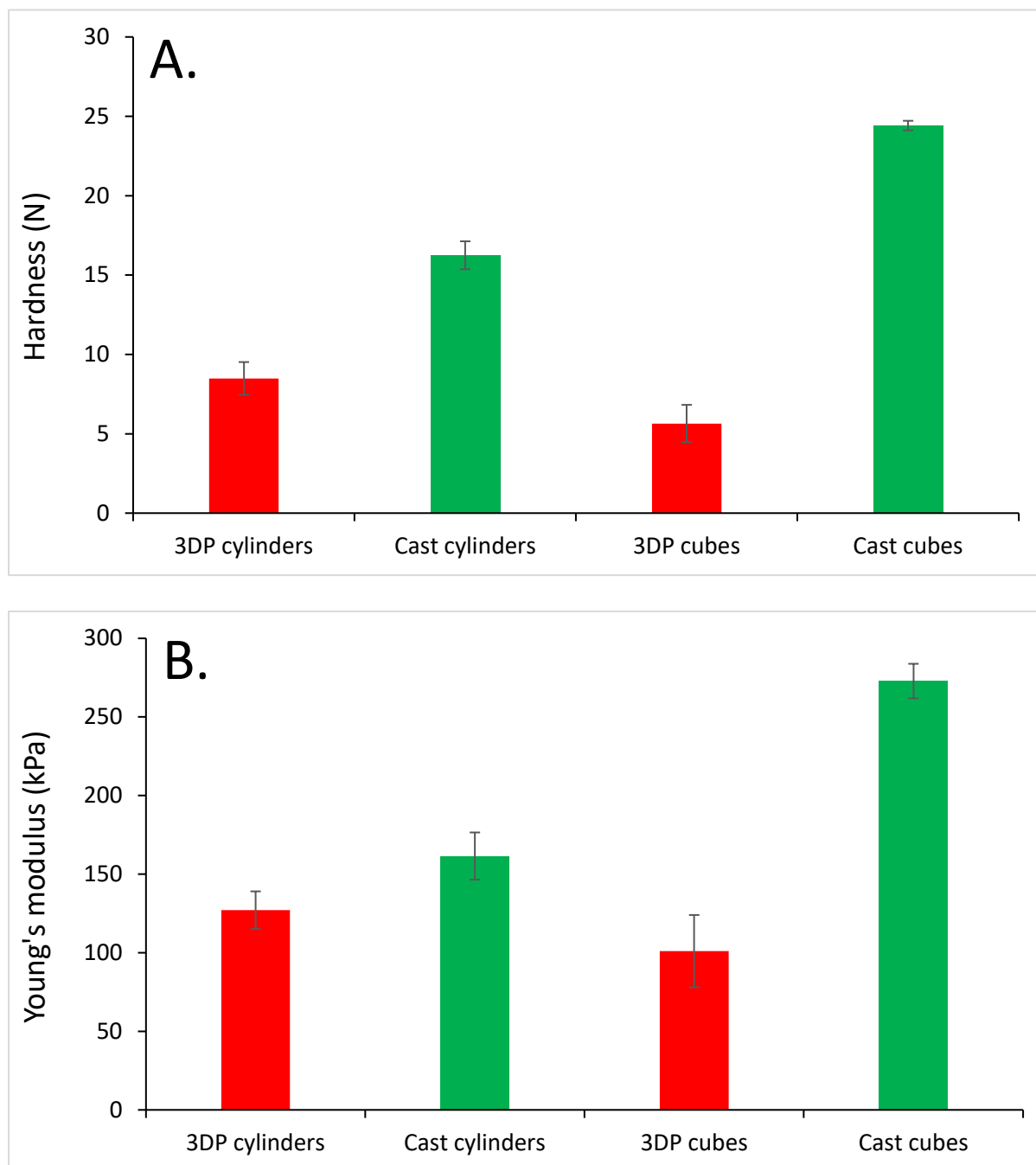


Figure 6: Hardness (A) and Young's modulus (B) of the printed and cast cubes and cylinders

The results obtained from the TPA are much higher than those observed in literature (Artignan, Corrieu, & Lacroix, 1997; David A Garrec, et al., 2013) which is due to both the higher concentration of  $\kappa$ C used and the addition of the  $\text{Na}^+$  ions and the thiamine reinforcing the gel network as discussed. The TPA data highlights the differences in the bulk structure of a printed gel compared to a cast gel. The higher hardness value shows that the continuous cast gel network is much more robust. The TPA showed that the cast samples were stiffer than the printed samples when undergoing compression. It was also noted that cubes were harder and less elastic than the cylinders for the cast samples. The cubes had a cross-sectional surface area of  $144 \text{ mm}^2$  and a volume of  $1728 \text{ mm}^3$ . The cylinders had a cross-sectional surface area of approximately  $113 \text{ mm}^2$  and a volume of  $1357 \text{ mm}^3$ . There exists almost no literature comparing cubes to cylinders of the same material subjected to

compression tests, within the range of materials studied, however it has been shown that variations in surface area can affect results obtained from TPA (Rosenthal, 2010). Since gels printed in this manner are in effect a discontinuous network, with several small networks only semi fused, the TPA showed they are less resistant to the external damage owing to the differences in the structure. Figure 7 shows a cube that has undergone compression testing. The printed shapes delaminated rather than fracturing like a cast gel. Since this was occurring rather than a fracture, the bonds holding the layers together must have been weaker than the gel network itself. Since the cast gels were one continuous network, they therefore could resist greater amounts of force as shown by the TPA results.

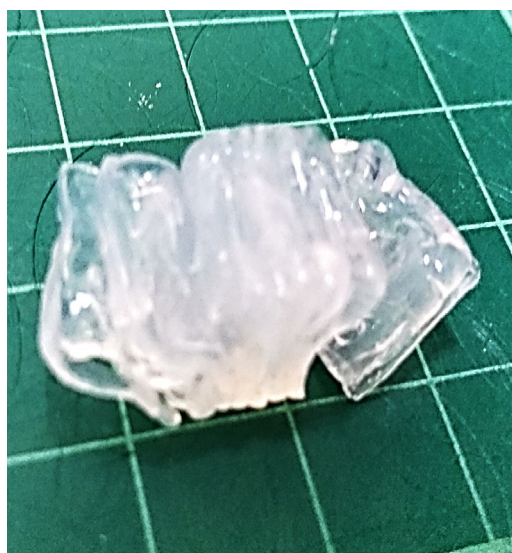


Figure 7: A printed 12 mm  $\kappa$ C-thiamine hydrogel cube that has delaminated after undergoing compression testing

### 3.4. Post-printing light microscopy of the hydrogels

While the separate printed layers could clearly be seen when observing the printed shapes with the naked eye, when cutting a central cross-section they were no longer discernible. It was important to establish whether the shapes had had some of the layers fuse together or whether the printed shapes were a series of individual gel networks held together by physical bonds. FDM printed plastics have been shown to have gaps running through the structure as a consequence of the manufacturing technique (Sood, Ohdar, & Mahapatra, 2010). The presence of these gaps running through the printed shapes could affect the final release profile of the thiamine by creating a shorter diffusion path. It also helped to confirm the findings from the TPA as the internal structure differed from that of a cast gel network. The differences in internal structure were responsible for overall different bulk structure. Figure 8A and B show the outside of a cast cube and a printed cube respectively. The ridges observed in 8B are due to the printing process and show a clearly different external structure to the cast cube. Figure 8C and D show a central cross-section from a cast cube and a printed cube. This highlights the stark differences in the structures created by 3DP and casting when it comes to hydrogel production. Figure 8D presented that 3DP produced hydrogels have visible layering running through them as indicated by the lines visible in the image. Figure 8C showed cast gels with a homogenous, continuous structure as was expected. This layering is believed to affect the physical characteristics of the gel as determined in the TPA.



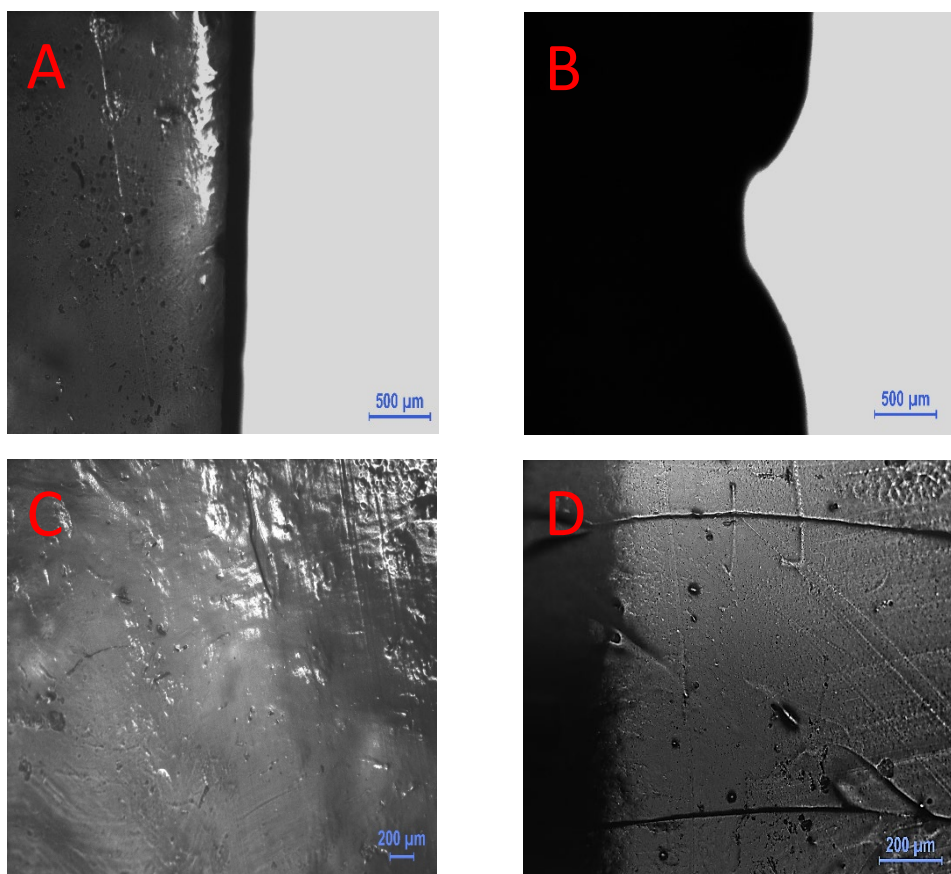


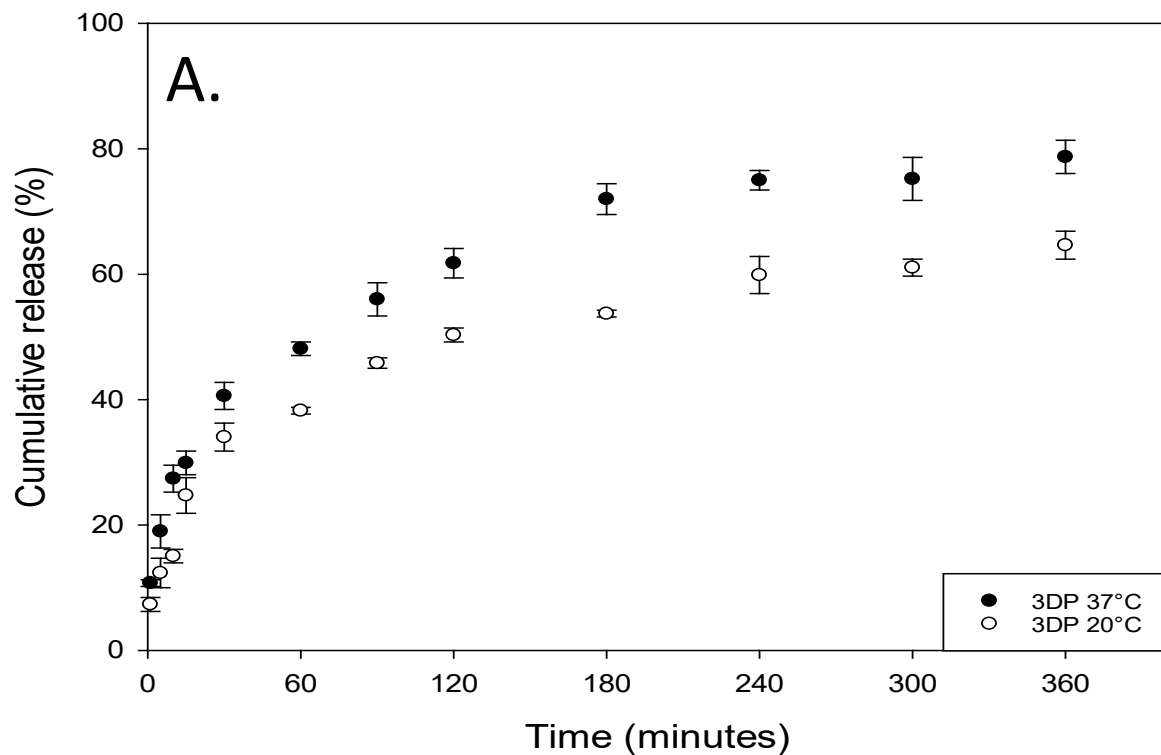
Figure 8: Microscope images of cast (A) and printed (B)  $\kappa$ C-thiamine hydrogels from the outside layer, and cross-sections of cast (C) and printed (D)  $\kappa$ C-thiamine hydrogels

### 3.5. Hydrogel release studies

There exists very little literature comparing the release rate from 3DP and traditionally manufactured structures. 3DP capsules have been shown to released dye at the same rate as injection moulded capsules (Melocchi, et al., 2015). However, these were only 0.3 mm in width and hollow, so this might not be applicable to the shapes studied which were far thicker and solid throughout. The amount of thiamine released from the printed and cast cylinders was assessed at 20°C and 37°C and the obtained release profiles are shown in Figures 9A and 9B respectively. Cubes were not selected for release testing as the chosen model for analysis does not account for this shape (Peppas, et al., 1989). All of the cylinders both printed and cast weighed  $1.45 \text{ g} \pm 5\%$ . Release of the active increased from 20°C to 37°C due to increased energy in the system facilitating a greater rate of release by diffusion (Vrentas & Vrentas, 1992). However, diffusion might not be the only factor in effect. Release of an active from a polymer matrix can also be dependent on relaxation of the polymer matrix as well as Fickian diffusion (Peppas, et al., 1989). The release data also showed a difference in the release rates between the printed and the cast gels. At both temperatures the printed gels showed an increase in initial release rate over the first fifteen minutes as shown by Figure 9C when compared to the cast gels. Moreover, the release data showed that there was a greater amount of thiamine released from the printed cylinders after 360 minutes compared to the cast ones.

Figures 9A and 9B showed that the cast cylinders released less thiamine over six hours at both of the tested temperatures compared to the 3D printed cylinders. Therefore, structural and mechanical differences between the two types of gel system must also contribute to the different release rates. In hydrogels, the physical characteristics determined from TPA such

as hardness have been shown to affect molecular release (Jones, Woolfson, Djokic, & Coulter, 1996). This was observed to occur in this study as well with the printed samples having lower values for hardness compared to the cast samples. Also, the elasticity seemed to have no effect on the release in their study with the elasticity values for the highest and lowest releasing systems being within 0.04 of each other on average. This indicates that while the TPA data might go some way in explaining the differences in release rate, it is not absolute. The layers running throughout the printed shapes lead to a decrease in the diffusion path as water is able to pass into the printed cylinders at a quicker rate than the cast cylinders owing to the differences in the bulk structure. This led to an increase in the release rate as shown by (Liang, et al., 2006). Furthermore, after the release tests the cast and printed cylinders had the surface water removed by drying and were weighed out, with the printed cylinders having increased on average more in weight than the cast cylinders. This further supports the notion that water was able to pass into the 3DP cylinders at a faster rate.



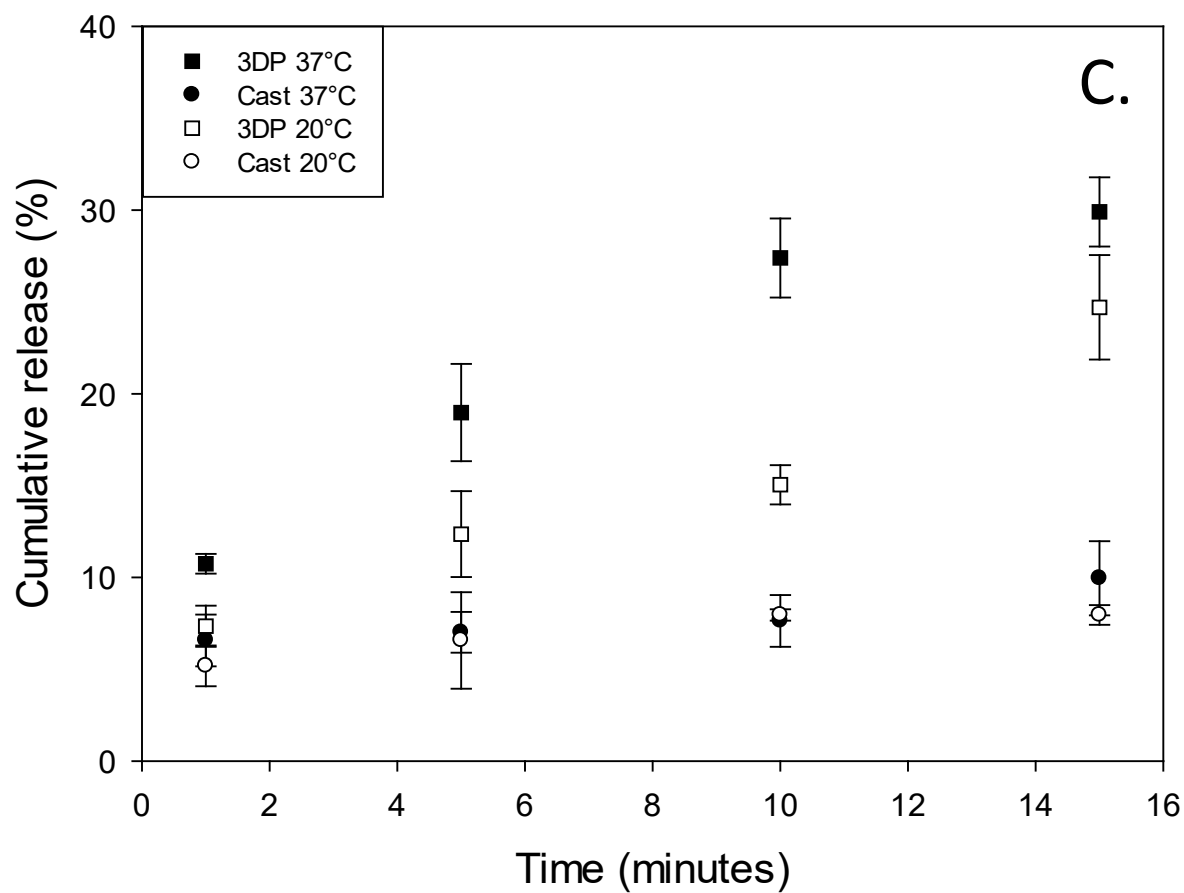
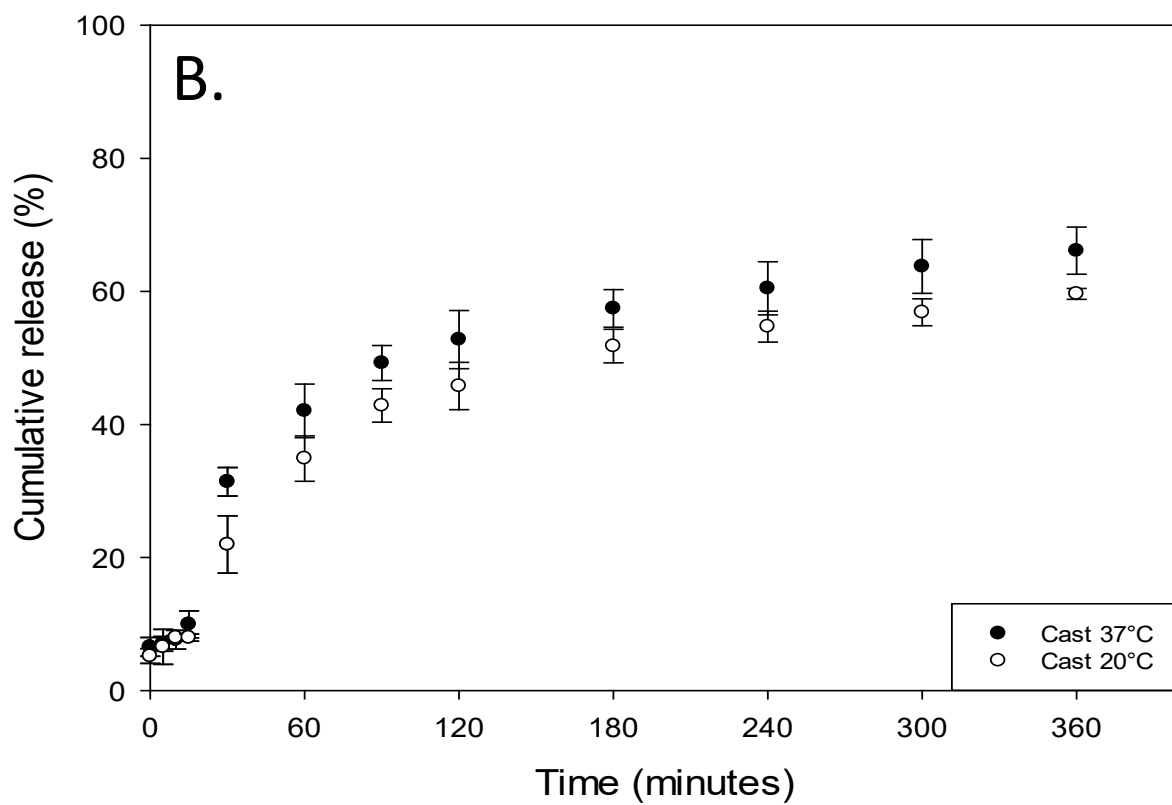


Figure 9: A comparison of cumulative release rates of thiamine from printed (A) and cast (B) and both for the first 15 minutes (C) κC 3% and thiamine 2% hydrogels.

The drug release profiles for the printed and cast gels at both temperatures were fitted to the Peppas-Sahlin equation. The kinetic constants for Fickian diffusion and case-II relaxational contribution were calculated through plotting the first 60% of the release data to equation 1. This made it possible to calculate F and R to the release of thiamine from the printed and cast hydrogels. The data is shown in Figure 10, with the information on  $k_1$ ,  $k_2$  and  $m$  available in table 2.

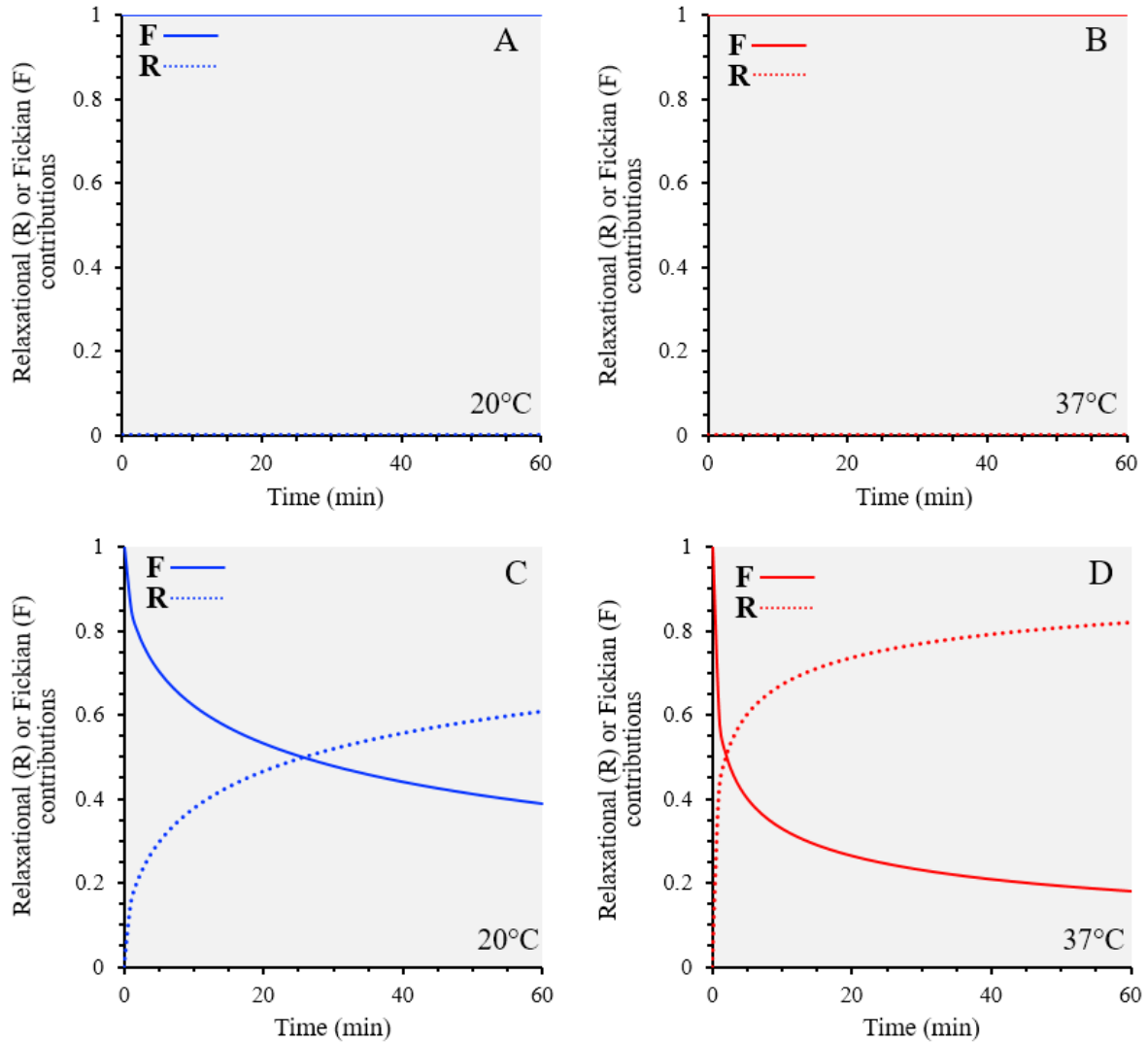


Figure 10: Data showing the percentage thiamine release due to Fickian diffusion and relaxation for printed cylinders at 37°C (A) and 20°C (B) and cast cylinders at 37°C (C) and 20°C (D)

	37°C printed	37°C cast	20°C printed	20°C cast
$k_1$	11.46	1.266	6.902	1.586
$k_2$	$8.37\text{E-}14$	0.9282	$2.91\text{E-}9$	0.2867
$m$	0.3572	0.4459	0.4321	0.5267

Table 2: The constant values used in the calculation for F and R for the release tests of the cast and printed cylinders at both temperatures

Figures 10A and 10B showed that with the 3DP shapes the diffusion contribution was essentially 100% for the printed shapes at both temperatures, with F being practically equal to 1 throughout the test. Whereas Figures 10C and 10D show that there is a relaxation contribution to the thiamine release from the cast cylinders. Both of the cast temperatures started out with diffusion being the dominant contribution. However, as the tests progressed the relaxation contribution started to exert more influence eventually becoming the dominant mechanism of release. This can be observed where the dotted line crosses the solid line. This happened more quickly at 37°C compared to 20°C due to there being more energy in the system. This led to a faster relaxation of the polymer chains (Watase, et al., 1968). This showed that the cast systems behaved in a manner seen in previous reports (Baggi & Kilaru, 2016) and (Lupo, Maestro, Gutiérrez, & González, 2015). However, for the 3DP cylinders it is believed that the different internal structure allowed water to penetrate faster into the shapes and so diffusion had been encouraged to such an extent as to make the relaxation contribution negligible (Falk, Garramone, & Shivkumar, 2004).

Figure 11A and B show the TPA data for the printed cylinders had a greater decrease in hardness and elasticity compared to the cast cylinders. By absorbing more water, essentially the concentration of the  $\kappa$ C and the thiamine within the gel decreased more compared to the cast gels. This caused there to be a less dense polymer network and could have contributed to an increase in release rate (Wu, Joseph, & Aluru, 2009). Also swelling would have increased the pore size of the hydrogel (Ganji, Vasheghani, & Vasheghani, 2010), which could also have increased the release rate in this case if the pore size was a rate limiting factor (Meena, Prasad, & Siddhanta, 2007). If it was not then the swelling would not have affected the release rate (Varghese, Chellappa, & Fathima, 2014). However, even after 48 hours, neither formulation had released 100% of the thiamine into the dissolution medium. This might be because with such a high  $G'$  value, the gel network was simply so dense that not all of the thiamine was able to diffuse out despite the swelling that occurred (Patil, Dordick, & Rethwisch, 1996). However, it could also be due to the use of an incubator shaker rather than the use of a USP paddle apparatus as is usually used in release studies. Finally, the electrostatic complexation of the  $\kappa$ C and the thiamine might also have been responsible for the failure to reach 100% release (Daniel-da-Silva, Ferreira, Gil, & Trindade, 2011). Future study needs to go into this area as well as studying different pH levels in the release media.

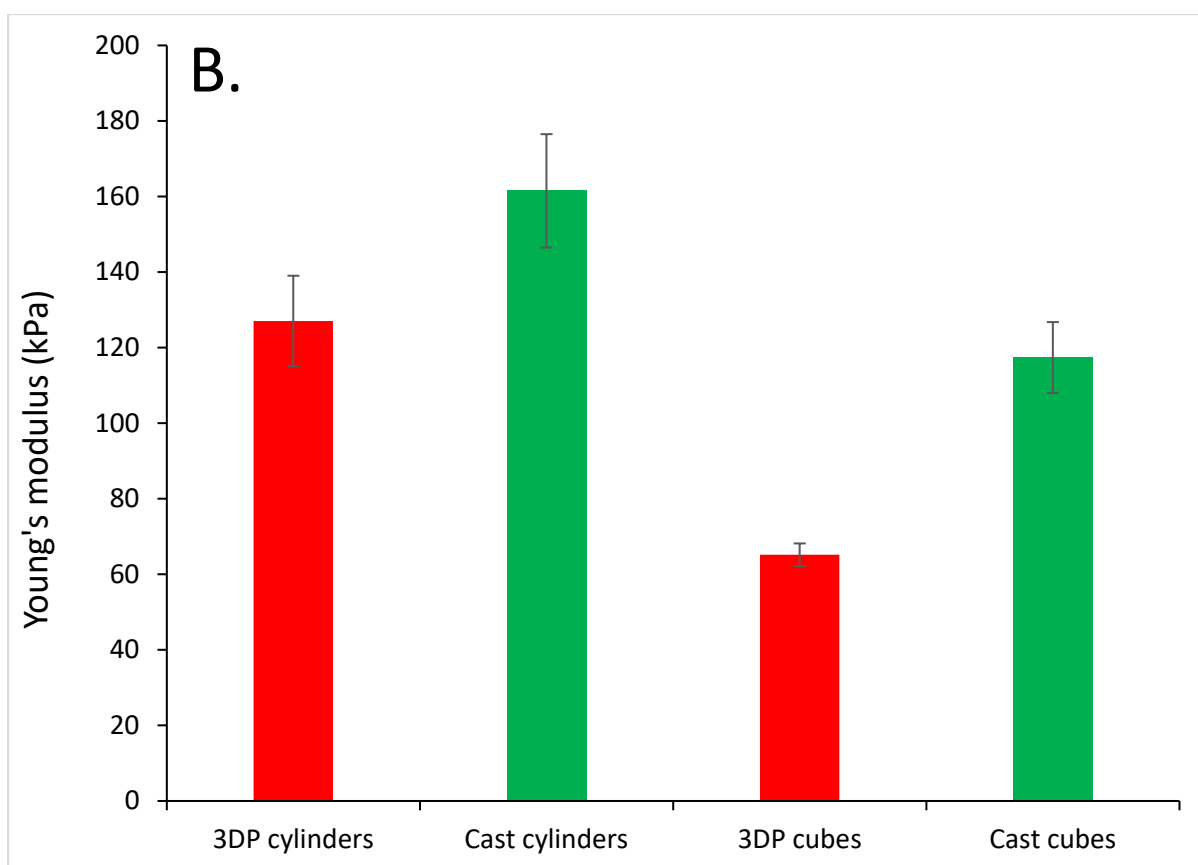
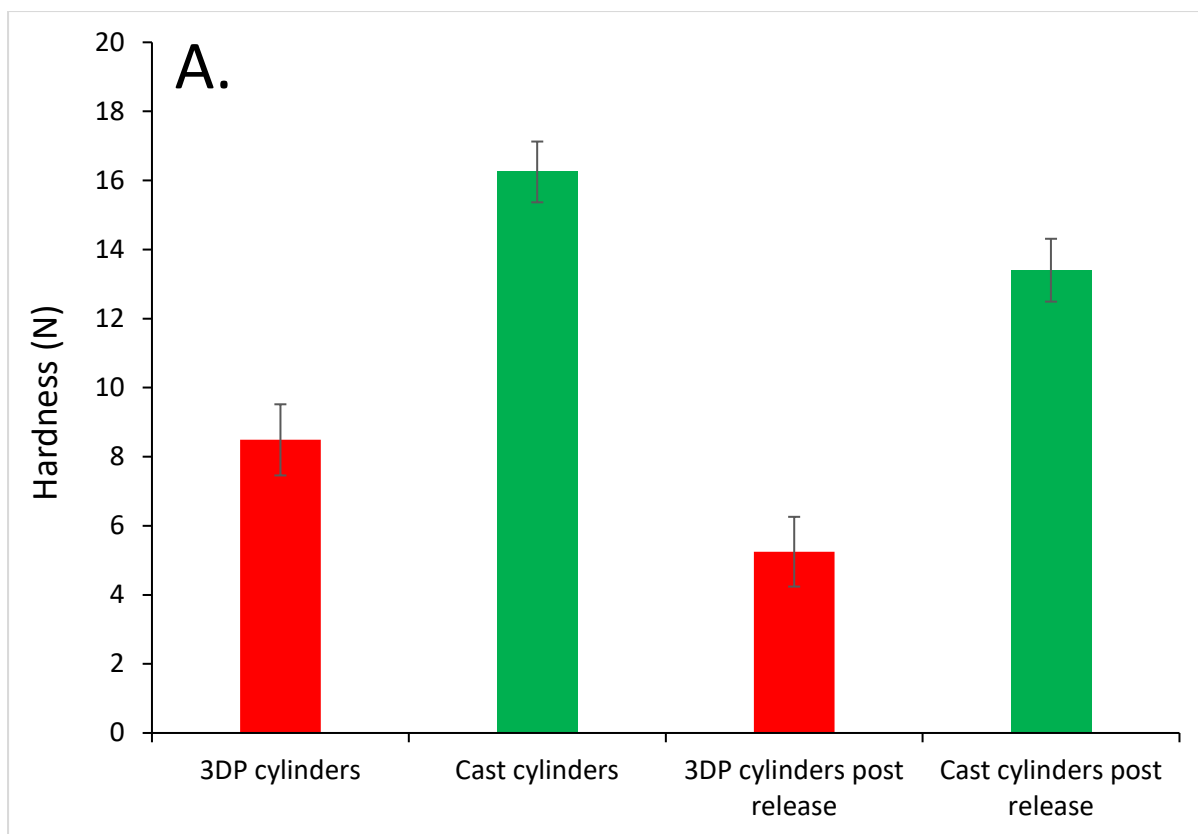


Figure 11: Post-release study texture analysis of printed and cast gels assessing hardness (A) and young's modulus (B)

## 4. Conclusions

In this study,  $\kappa$ C and agar thiamine-loaded hydrogels were assessed for their suitability for 3D printing. Rheological analysis showed that while thiamine does not interact with agar, it can form electrostatic complexes with  $\kappa$ C that lead to gel structure reinforcement (up to a point). Beyond this, thiamine addition caused a marked decrease in  $G'$  while both  $T_{gel}$  and  $T_{melt}$  continued to increase.  $\mu$ DSC confirmed the same trends and further revealed that the gelling and melting enthalpies for  $\kappa$ C-thiamine systems were shown to decline with increasing thiamine concentration.

2% agar was unable to print due to a lower  $G'$  and  $T_{gel}$ , but 3%  $\kappa$ C-2% thiamine hydrogels were printable, and cubes and cylinders could be formed with reproducible weights and dimensions. The printed gels' physical properties were then compared to traditionally produced cast gels of the same dimensions and weight within a margin of  $\pm 5\%$ . TPA, microscopy and release studies were used to show that 3D printing facilitated the creation of gels with different physical properties without having to chemically modify the gel ingredients.

In terms of release, printed cylinders released a higher fraction of enclosed active than the cast cylinders. They also exhibited a faster rate of release over the first 15 minutes of testing. This was due to differences in the physical structure of the printed cylinders compared to the cast cylinders, which meant that printed cylinders were more prone to swelling. Modelling showed that while thiamine release from the cast cylinders was driven by both Fickian and relaxation phenomena, printed cylinders allowed the delivery of the active solely via diffusion.

Although agar was not shown to be suitable for printing under the current formulation/processing conditions, printability might still be realised at higher hydrocolloid concentrations or through the use of adjunctive materials to increase its gelation rate and  $G'$  value. Similarly, although  $\kappa$ C hydrogels could be successfully printed, they were unable to release the entirety of enclosed thiamine, thus further flexibility in terms of active delivery could be achieved for non-interacting addenda. Nonetheless, the current study does clearly highlight the promise of hydrogel utility for formation of food-related structures via 3D-printing, capable of performing differently to cast gels of the same material.

## Acknowledgements

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