Layered cryogels laden with Brazilian honey intended for wound care

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Abstract

PVA cryogels are well established as candidate biomaterials for wound healing applications but are not themselves biodegradable or antimicrobial. Blending PVA with NaCMC (CMC) or gelatin (G) can increase the gel’s ability to swell and would introduce a degree of biodegradability. The incorporation of appropriate amounts of a natural antimicrobial/healing agents, such as honey (H), would contribute to the gels properties. The present work addresses the development and characterization of layered gels (PVA-H, PVA-CMC-H and PVA-G-H, with empty PVA, PVA-CMC, PVA-G gels presented as controls). The gels were characterized by FTIR, DSC, in vitro analysis of swelling and microbiological (S. aureus) effects. Addition of gelatin, NaCMC and honey to PVA diminished the PVA chains’ ability to pack into crystallites. Samples containing honey swelled less and presented higher weight loss/biodegradability than samples without honey. Only the honey-laden PVA-CMC and PVA-G presented activity against S. aureus.

Keywords: layered hydrogel, PVA, NaCMC, gelatin, honey.

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1. Introduction

Infection associated with wounds to the skin affect 14 million people per year in the USA. Such infections are characterized by colonization by gram-positive bacteria such as S. aureus in the early stages of healing, which are later replaced by gram-negative organisms[1]. Requirements for successful healing include a mechanically stable and moist environment, a capacity for absorbing wound exudate, and antimicrobial properties which act against the development of infection.

Hydrogels have been manufactured in layers to mimic the layers and function of the skin. For example, chitosan-g-poly(ethylene glycol) hydrogel reinforced with chitosan-alginate were designed to mimic the micro-environment relevant to skin tissue engineering[6]. A hydrogel composed of layers of alginate/chitosan/poly(-glutamic acid) increased wound epithelialization and collagen regeneration[3]. Layered hydrogels based on PVA (polyvinyl alcohol) have also been developed: 2-layered skin equivalent PVA or PVA-cellulose fibre blocks were prepared by freeze-thawing and presented roughness and elasticity similar to human skin[4]. A PAA layer has been added to a PVA layer, with the PAA solution poured on the top of PVA swollen layer, where the chains of both polymers would entangle and form hydrogen bonding at the interface. The presence of PAA led to increased swelling, adhesion and biocompatibility while the presence of PVA underpinned the mechanical properties of the gel[5]. Hydrogels are potential materials for burn dressings.
PVA is a hydrophilic biocompatible polymer used to manufacture hydrogels. PVA hydrogels are capable of presenting physical properties similar to human tissue, such as elasticity[6]. Hydrogels can be based on chemical crosslinking or physical crosslinking, where chemical crosslinking is based on the use of crosslinking agents, such as glutaraldehyde, or based on the use of radiation[7,8]. Physically crosslinked PVA hydrogels can be prepared by freezing and then thawing an aqueous solution of PVA. PVA chains entangle when in solution. When frozen, ice crystals are formed. By phase separation (ice/PVA), PVA chains are pushed together forming crystallites, which are responsible for the high mechanical properties of these gels. These gels have tissue-like elasticity, toughness and are non-toxic[9].

Blended PVA gels incorporating several additives, such as L. bulgaricus extract[10] and neomycin sulfate[11], have been studied in the literature with respect to antibacterial and healing properties.

Sodium carboxymethyl cellulose (NaCMC) is a hydrophilic polymer, a polyelectrolyte, derived from cellulose. The charges of this polyelectrolyte elongate the polymer chains, increasing the water uptake capability of the network, with counterions favouring water entrance[12]. NaCMC is able to form hydrogels by blending it with hydroxyethyl cellulose and chemically crosslinking with divinylsulphone[13]; NaCMC can be combined with protein sericin, crosslinked by freeze-thawing, with glutaraldehyde and AlCl₃ to produce hydrogels for wound dressing[14]. NaCMC can also be blended with polyethylene glycol and crosslinked with citric acid for dressing purposes, showing no cytotoxicity[14]. In order to present antibacterial properties, NaCMC hydrogels can be loaded with drugs. For example, NaCMC gels can be crosslinked with citric acid and loaded with MCM-41 mesoporous silica nanoparticles containing tetracycline (antibiotic), where higher particle content led to higher S. aureus inhibition[15]. In another study, NaCMC was blended to PVA to physically (freeze-thawing) form hydrogels loaded with sodium fluoride, where the addition of NaCMC increased the gels swelling capacity, vapor transmission rate and porosity[16]. PVA-NaCMC-sodium fluoride hydrogel showed faster healing compared to a PVA-NaCMC gel[17].

Gelatin is a water-soluble protein obtained from the thermal denaturation of collagen. Gelatins with high levels of amino acids present high gel strength and melting point[18]. Gelatin is biocompatible, biodegradable, it has the ability to form films, and is generally low cost. In order to avoid dissolution in aqueous fluids, gelatin hydrogels have to be crosslinked with glutaraldehyde, genipin or lactose. Gelatin hydrogels crosslinked with lactose have been investigated as wound dressing materials[19]. Chitosan/gelatin crosslinked hydrogels were loaded with phenols from Hamamelis virginiana and showed activity against P. aeruginosa and S. aureus[20]. Superabsorbent gels were prepared by esterification of PVA with gelatin[21]. Chitosan/gelatin/PVA irradiated hydrogels for wound dressing applications presented swelling capacity, adequate vapor transmission rate and mechanical properties[22]. PVA/chitosan/gelatin hydrogels loaded with PCL microspheres containing bFGF and BSA (protein) showed no cytotoxicity, and an improved healing rate, while chitosan improved the antimicrobial properties[23].

Honey is a bee-based product which can present more than 200 constituent components, mainly carbohydrates (most of them monosaccharides, e.g. fructose, glucose, sucrose), proteins, organic acids, vitamins and phenolic compounds. Honey’s antimicrobial characteristics are related to the H₂O₂ produced by enzymatic activity and to the action of complex phenols and organic acids (flavonoids)[24-26]. The honey composition is influenced by the flower types visited by bees, the soil composition, bee species and climatic conditions. Brazilian honey presents a considerable variability, since the local flora and climate change all over the country. The honey’s consistency, color, odor, taste and scent are constant physical-chemical characteristics in monoflower honeys. Honey’s phenolic substances and flavonoids are responsible by pharmacological properties, e.g. antimicrobial and antioxidant activities[26]. Wounded rats treated with Brazilian honey (from southwest region of Brazil) presented fastest recovery than the control group, it also treated the wounds infection[27]. Brazilian honey, also from the southwest, was tested against S. aureus microorganisms and it showed antimicrobial activity, due to its phenolic compounds[28]. Brazilian honey shows potential to be used as wound healing agent.

Honey has been successfully incorporated in hydrogels. For example, a chitosan dressing containing 75% honey enhanced tissue regeneration[29]. A pectin-honey hydrogel promoted faster wound healing than a control[30]. PVA crosslinked with borax and loaded with honey promoted the proliferation of cells, with swelling, and permeability properties considered adequate for moderate exudative wounds[31]. Gelatin-chitosan hydrogels loaded with compounds from manuka honey presented antibacterial activity, a high wound healing rate and the absorption of exudates[32]. Gelatin-chitosan-honey hydrogel was effective against S. aureus and E. coli and stimulated burns healing[33]. PVP-agar-Peg gels containing 6% of honey accelerated rat wounds contraction[34]. PVA/chitosan/montmorillonite gels loaded with 15% Iranian honey showed potential to absorb exudate and efficient wound healing[35]; acrylamide hydrogels were loaded with different amounts of Indian honey (5-15%), but the gel containing 10% honey was considered the best one, due to optimized tensile strength and swelling capacity[36]; carbopol 934 and chitosan gels containing 75% Egyptian honey presented high burning healing rate[37].

The goal of the present work was to develop layered PVA, PVA-honey (PVA-H), PVA-NaCMC (PVA-CMC), PVA-NaCMC-honey (PVA-CMC-H), PVA-gelatin (PVA-G), PVA-gelatin-honey (PVA-G-H) hydrogels and characterise their physico-chemical and functional properties in order to gain insights into their potential development as wound-care biomaterials.

2. Materials and Methods

2.1 Sample preparation

The samples were manufactured by the dissolution of PVA (Sigma-Aldrich, Mw: 85,000-124,000 Da, 99.9% hydrolyzed) at 90 °C under mechanical stirring for 4h according to Table 1. For each PVA blend without honey, the amount of water used was split in half to separately dissolve...
Table 1. Samples composition.

| Sample                | PVA  | NaCMC | gelatin | honey | H2O   |
|-----------------------|------|-------|---------|-------|-------|
| PVA (3 layers)        | 10g  | -     | -       | -     | 100 mL|
| PVA-CMC (3 layers)    | 8g   | 2g    | -       | -     | 100 mL|
| PVA-G (3 layers)      | 8g   | -     | 2g      | -     | 100 mL|
| PVA-H – 1st layer     | 10g  | -     | -       | -     | 100 mL|
| PVA-H – 2nd layer     | 10g  | -     | -       | 10 mL | 90 mL |
| PVA-H – 3rd layer     | 10g  | -     | -       | 5 mL  | 95 mL |
| PVA-CMC-H – 1st layer | 8g   | 2g    | -       | -     | 100 mL|
| PVA-CMC-H – 2nd layer | 8g   | 2g    | -       | 10 mL | 90 mL |
| PVA-CMC-H – 3rd layer | 8g   | 2g    | -       | 5 mL  | 95 mL |
| PVA-G-H – 1st layer   | 8g   | -     | 2g      | -     | 100 mL|
| PVA-G-H – 2nd layer   | 8g   | -     | 2g      | 10 mL | 90 mL |
| PVA-G-H – 3rd layer   | 8g   | -     | 2g      | 5 mL  | 95 mL |

The PVA and either NaCMC (Sigma-Aldrich, average Mw ~250,000, degree of substitution 0.9) or Gelatin (from bovine skin, Sigma-Aldrich, Type B). The resulting solutions were then mixed together after each solution had reached room temperature (under stirring). The NaCMC was dissolved in water under stirring at room temperature for 2h. Gelatin was dissolved at 60 °C under stirring for 4h.

The samples were prepared in three steps using 24-well plates. The composition of each layer of the samples without honey was similar (PVA or PVA-CMC or PVA-G). The first layer was obtained by pouring 1 mL of the polymer solution or the blend solution per well followed by a freeze-thaw cycle (each freeze-thawing cycle was to submit the sample to 1h at -18 °C and 30 min at room temperature). The second layer (1 mL/well) was subsequently added and freeze-thawed, followed by the addition and freeze-thawing of the third layer. The first layer was then subjected to 3 freeze-thawing cycles, the second layer to 2 cycles and the third layer to 1 cycle.

For the samples that were to contain honey (a commercial Brazilian honey from Southeast region, characterized as “silvestre” honey), the polymers were dissolved in the designated amounts of water from Table 1 following the procedures previously described. Honey was then added to the polymer solution or to the blend solution, according to the quantities indicated in Table 1, at room temperature under mechanical stirring for 5 minutes. The first layer of these samples was either pure PVA or a PVA based polymer blend without honey and had experienced 3 freeze-thawing cycles. The second layer was the polymer/blend with 10% honey, submitted to 2 freeze-thawing cycles; and the third layer was polymer/blend with 5% honey, submitted to 1 freeze-thawing cycle. All the samples were dried in oven at 50 °C for 24h. Regarding honey’s thermal degradation, honey samples treated at 23 °C for short times did not degrade, but the same behavior was not observed for samples heated at 95 °C, although antioxidant activity might increase with the heating temperature[28]. Heat treated honey might present hydroxymethylfurfural, which should be lower than 80 mg/kg in tropical honeys, since it is toxic and carcinogenic. In addition, tropical honeys can present fermentation when heat treated, but it can be prevented by heating honey at 60-70 °C for 10 min or at 60-65 °C for 30 min[29]. The drying temperature was low to preserve honey activities, although honey samples autoclaved at 121 °C could maintain its properties[30].

2.2 Physico-chemical analysis

The physico-chemical analysis of the samples was performed by Fourier Transformed Infrared Spectroscopy (FTIR), equipment VERTEX-70 (UFRJ), 32 scans per sample, from 4000 cm-1 to 600 cm-1. Samples representing each layer were prepared separately and submitted to the proper number of freeze-thawing cycles separately and then dried to be evaluated by FTIR. The multi-layered samples were also evaluated by FTIR.

2.3 Thermal analysis

The samples were analysed by Differential Scanning Calorimetry (DSC), where approximately 5 mg of each sample composition was submitted to 10 °C/min of heat flow from room temperature to 240 °C using a DSC Q200 TA Instruments equipment (EMBRAPA Agroindustry). The transition temperatures of the samples (glass transition temperature - Tg, melting temperature - Tm) were obtained using the second cycle of heating to overcome the samples’ thermal history. The samples degree of crystallinity (Xc) was calculated by

\[
X_c = \frac{100\Delta H_m}{\Delta H_{m100}}
\]

The \(\Delta H_m\) is the enthalpy of melting related to the samples at the PVA melting temperature and \(\Delta H_{m100}\) is the enthalpy of melting related to PVA 100% crystalline, 138.6 J/g[26].

2.4 In vitro analysis

The swelling tests proceeded in saline solution at room temperature. For each sample composition, 5 samples were cut (samples weight was standardized among the ones with the same composition) and immersed in saline solution (20 mL per sample, used to mimic the body fluids). The samples were removed from the saline solution at regular intervals (30 min, 1h, 2h, 3h, 4h, 24h, 48h, 72h and 96h). The adsorbed fluid was removed using filter paper and the samples were weighed and then returned to the media. The swelling degree (SD) was calculated according to

\[
SD = \frac{100(W_S - W_{DB})}{W_{DB}}
\]
W_{db} is the weight of dried samples before swelling; W_{s} is the weight of the swelled samples.

At the end of the 4 days of immersion, the samples were dried in oven at 50 °C for 24h and weighed to calculate the sample gel fraction (GF)

\[ GF = \frac{100W_D}{W_{db}} \]  

(3)

W_{D} is the weight of the dried samples after swelling) and weight loss (WL)/biodegradability\[34],

\[ WL = 100\left(\frac{W_{db} - W_D}{W_{db}}\right) \]  

(4)

For microbiological analysis, the ASTM E2180-07 standard was adapted. A suspension of *Staphylococcus aureus* cells (ATCC 6538) was prepared (guaranteeing that the suspension reached 0.5 in the MacFarland scale), which correspond to 10⁶ colony forming unit (UFC)/mL. The agar paste received 1 mL of the mentioned suspension, where the agar paste total amount of microorganisms were 10⁶ UFC/mL. Each sample was placed in an empty well of 24 wells flat bottom polystyrene plate, and each well received 200 μL of the agar paste containing *S. aureus* (where duplicates for each samples composition were evaluated). The plates were incubated at 30 °C for 24h. After incubation, the media in contact with samples were placed in Falcon tubes. After that, different amounts of buffer solution were added to prepare decimal dilutions of the media in contact with samples and to subsequently, count the colony forming units (cfu) by the micro dropping technique. The bacterial calculation was performed using the optical microscope (equipment Olympus). The reduction calculation was based on ASTM Standard 20170504.

### 3. Results and Discussions

It was not possible to distinguish the samples layers, indicating that the system might behave homogeneously. All samples, containing honey or not, presented homogeneous morphology, similar to PVA-H, Figure 1.

Figure 1. PVA-H sample section.

3.1 Physico-chemical analysis

The FTIR spectra of all samples are shown in Figure 2. Each layer of PVA presented similar FTIR spectra of the three-layered sample. Nonetheless, the band at 1142 cm⁻¹, related to the formation of PVA crystallites (intra- and intermolecular hydrogen bonding between the chains originated by hydrophilic forces)\[35], presented lower transmittance with increasing numbers of freeze-thawing cycles, indicating the contribution of the freeze-thawing process to the formation of PVA crystals. There were (in all PVA layers) bands at: 3273 cm⁻¹ (OH hydrogen bonds)\[36]; 2941 cm⁻¹ and 2909 cm⁻¹, \(\nu(C-H)\) alkyl\[37]; 1652 cm⁻¹, \(\nu(C=O)\), from residual aldehyde\[38,39]; 1563 cm⁻¹ and 1237 cm⁻¹, \(\nu(C=O)\)\[35]; 1411 cm⁻¹, \(\delta(-CH)\)\[37]; 1380 cm⁻¹, \(\omega(C-H)\)\[38]; 1329 cm⁻¹, \(\delta(CH=O)\)\[35]; 1089 cm⁻¹, \(\nu(C-O)\)\[38]; 916 cm⁻¹, \(\delta(-CH)\)\[39]; and 835 cm⁻¹, \(\nu(C=O)\)\[39].

The previously reported PVA bands were also observed in the PVA-CMC as well as the overlapping bands of NaCMC, showing the dispersion of NaCMC in the PVA matrix\[40]; a shoulder at 2855 cm⁻¹, \(\nu(C-H)\)\[41]; 1585 cm⁻¹ (non-hydrated \(\nu(C=O)\) group)\[42]; 1415 cm⁻¹, \(\nu(COO-)\); 1325 cm⁻¹, \(\delta(C-H)\) of methyl groups; 1080 cm⁻¹ \(\nu(C-O)\)\[43]. The band at 1652 cm⁻¹ (\(\nu(C=O)\) is absent in the sample PVA-CMC\[35]. The band at 1560 cm⁻¹ (carbonyl group) in the PVA-CMC sample has a different shape compared to the same band in the PVA sample, indicating a change in the balance of free associated carbonyl groups. It could relate to the polymers miscibility, although displacement of the bands related to the crystallinity of PVA\(1142 \text{ cm}^{-1}, \nu(C-C)\) and of NaCMC\(1372 \text{ cm}^{-1}, \delta(CH)\) and displacement of the NaCMC \(\beta\)-glucosidic groups were not identified. As expected, the PVA crystallinity index (ratio between the absorbance of the band at 1142 cm⁻¹ and the band at 2905 cm⁻¹) is lower in the blend, \(I_{\text{PVA}} = 1.45\) and \(I_{\text{PVA-CMC}} = 1.36\), where the presence of NaCMC diminishes the PVA chains ability to pack into crystals\[44]. The PVA-CMC layers revealed that the increase of freeze-thawing cycles displaced the band at 1593 cm⁻¹ (NaCMC’s \(\nu(-COO^-)\)) to 1566 cm⁻¹, which indicate the effectiveness of the “crosslinking”\[45]. In addition, the two bands at 1087 cm⁻¹ (PVA’s \(\delta(C-O-H)\)\[46]) and at 1060 cm⁻¹ (NaCMC’s \(\nu(OCH–O–CH)\)\[47]) cannot be distinguished in layers submitted to more freeze-thawing cycles, where only one band at 1087 cm⁻¹ can be observed.

The PVA-G sample presented some differences when compared to the PVA sample, e.g. the band at 1646 cm⁻¹ was more intense in the PVA-G sample, which could be due to the gelatin \(\nu(C=O)\)\[38]. There was a band at 1169 cm⁻¹ in the PVA-G sample (absent in the PVA sample); the PVA bands at 1142 cm⁻¹ and at 1089 cm⁻¹ were displaced to 1136 cm⁻¹ and to 1101 cm⁻¹ in the PVA-G sample, respectively. The gelatin bands displacement towards higher wavenumbers could be related to reaction products, e.g. from 1680 cm⁻¹ to 1758 cm⁻¹ would represent the formation of esterified product\[47]. There were bands at 640 cm⁻¹ and at 612 cm⁻¹ in the PVA-G sample, while there was no band in this region in the PVA sample. A band at 670 cm⁻¹ would be attributed to the gelatin \(\gamma(N-H)\), although bands observed at lower wavelengths were not reported to gelatin\[38]. The PVA-G layers revealed similar bands to those of the whole PVA-G sample.
The PV A-H sample presents bands related to PV A, e.g. at 3275 cm\(^{-1}\), 2940 cm\(^{-1}\), 1564 cm\(^{-1}\), 1379 cm\(^{-1}\), 1239 cm\(^{-1}\), 1143 cm\(^{-1}\). Nonetheless, some of its bands can be related to the presence of honey in the samples, e.g. at 1646 cm\(^{-1}\) (ν(C–H) of carboxylic acids, ν(NH\(_3\)) of free amino acids, water δ(OH)[48]), 1417 cm\(^{-1}\) (δ(O–H) of the C–OH group and δ(C–H) of the alkenes[48]), 775 cm\(^{-1}\) (saccharide configuration; anomeric region of carbohydrates vibration or δ(C–H)[49]). In addition, some of the honey bands are slightly displaced in the PV A-H sample, e.g. the bands at 1054 cm\(^{-1}\) (ν(C–O) of the C–OH group and carbohydrate structure’s ν(C–C)[48]); 1030 cm\(^{-1}\) (vibration of the C–OH group, carbohydrate structure’s ν(C–C) and ν(C-O), phenol’s C-O vibration[50]); 898 cm\(^{-1}\); 874 cm\(^{-1}\), displaced by ~9 cm\(^{-1}\); 819 cm\(^{-1}\), these last bands related to saccharide configuration and to anomeric region of carbohydrates vibration or δ(C–H)[49]. Some of the PV A-H bands are located in between PV A bands position and honey bands position, e.g. the band at 1336 cm\(^{-1}\), which is in between the honey band at 1345 cm\(^{-1}\) (flavano1’s and phenol’s δ(O-H), δ(C-O), δ(C-H) and δ(C=C)[50] and the PVA band at 1329 cm\(^{-1}\)[35]); there is a shoulder at 1097 cm\(^{-1}\), between the PVA band at 1089 cm\(^{-1}\) (ν(C=O)[39]) and the honey band at 1100 cm\(^{-1}\) (ν(C=O) related to the C=O–C linkage[48]); 917 cm\(^{-1}\), which is between the PVA band at 916 cm\(^{-1}\)[39] and the honey band at 918 cm\(^{-1}\) (carbohydrate’s δ(C–H)[49]; α and β anomers’ ν\(_a\)(C-O-C)[53]). These bands displacement and the bands located in between the original PVA and honey bands could indicate physical interaction (Van der Waals or hydrogen bonding) between PVA and honey.

Since the layers of the samples presented varied composition, they were examined separately. The layer containing 5% honey presented bands similar to the PVA-H whole sample previously described, but the bands at 1336 cm\(^{-1}\) and at 898 cm\(^{-1}\) (honey’s bands)[52] were absent and a shoulder at 831 cm\(^{-1}\) (PVA’s band)[39] was present; the band that should be at 1054-1030 cm\(^{-1}\) was displaced towards the PVA band (1089 cm\(^{-1}\)[39]), it was at 1083 cm\(^{-1}\).

The layer containing 10% honey also present the same bands as the PVA-H whole sample, but the bands at 2940 cm\(^{-1}\) and at 2910 cm\(^{-1}\) were absent. Nonetheless, there were bands at 2930 cm\(^{-1}\) (honey’s band[48]), 1338 cm\(^{-1}\), 1259 cm\(^{-1}\) (honey’s band[50]), 1189 cm\(^{-1}\) (honey’s band[50]). A slight displacement of PVA and of honey bands in the PVA-H layers could be observed with the increased honey content, which could indicate the effect of honey in the movement of PVA’s functional groups and vice versa. The increased level of honey in the layer also revealed more honey bands in the FTIR spectrum, as expected.

The PVA-CMC-H sample presented the same bands as the PVA-CMC and that were previously described. In addition, there were bands related to honey, e.g. 2923 cm\(^{-1}\), 1417 cm\(^{-1}\), 1056 cm\(^{-1}\), 1035 cm\(^{-1}\) and 819 cm\(^{-1}\). These bands are slightly displaced in the PVA-CMC-H sample. The layer containing 10% honey and the layer with 5%
honey presented similar bands to PVA-CMC-H whole sample previously described, although there were slight displacements of the bands with the addition of honey, e.g. the band at 1088 cm$^{-1}$ in PVA-CMC layer is broadened and shifted to 1074 cm$^{-1}$ in the layer containing 10% honey. There is a band at 1592 cm$^{-1}$ in the PVA-CMC-H layers that is absent in the PVA-CMC layer and in the honey spectrum. The presence of a new band could indicate new bonding between the polymers and honey\cite{53}.

The PVA-G-H sample presented the previously described PVA bands; some gelatin bands, e.g. at 1551 cm$^{-1}$ and at 1337 cm$^{-1}$\cite{54}; and some honey bands previously described, e.g. at 1052, 1029, 896, 853, 817, 775 cm$^{-1}$. Band displacement or the formation of bands related to a chemical reaction or interaction between honey and the polymers was not observed\cite{21}.

### 3.2 Thermal analysis

DSC analysis of the whole three-layer samples revealed that the addition of the polysaccharide (NaCMC) diminished the sample $T_s$ and raised the sample $T_m$, compared to PVA, while $X_c$(PVA-CMC) was lower than the $X_c$(PVA) (Table 2). It suggests that the NaCMC chains do not contribute to polymer chain entanglement, probably working as a plasticizer in the amorphous region of the blend. In addition, the NaCMC diminished the degree of crystallinity, but more “perfect” crystals were formed (with a higher $T_m$). A similar effect was observed when gelatin was added to PVA, but the $T_m$ of PVA and PVA-G was similar, indicating that gelatin contributed only as a plasticizer and to diminish the sample’s degree of crystallinity. The addition of honey to PVA diminished considerably the $T_s$, $T_m$ and the $X_c$ of the samples, indicating that honey is a physical barrier to the interaction of polymer chains. The addition of both honey and either polysaccharide or protein to PVA led to the absence of crystallinity (there is no peak related to the crystalline phase). The PV A samples weight loss was similar to PVA-CMC-H, but the $T_m$ of PV A samples was lower than the $T_m$ of PVA-CMC-H layers, indicating that the samples crystallites work according with the decrease in the samples’ degree of crystallinity. Samples that are less crystalline present more amorphous phase and thereby stretch more available to the saline solution\cite{59}. PV A samples swelled less than PVA-G samples (which have a low degree of crystallinity compared to PVA), and these in turn swelled less than PVA-CMC samples (which have an even lower degree of crystallinity). Samples that are less crystalline present more amorphous phase and thereby stretch more available to the saline solution. PV A-CMC-H layers can be considered the most similar sample in terms of swelling degree (SD) between all the samples, although the SD of the samples PV A and PV A-CMC-H can be considered the same. In general, it can be observed that samples containing honey swelled less than samples without honey\cite{58}, probably because honey occupies network pores created by the ice during freeze-thawing, filling space that could have been available to the saline solution\cite{59}. PV A samples swelled less than PVA-G samples (which have a low degree of crystallinity compared to PVA), and these in turn swelled less than PVA-CMC samples (which have an even lower degree of crystallinity). Samples that are less crystalline present more amorphous phase and thereby stretch more fluid ingress, and therefore swell more\cite{60,61}.

The gel fraction (GF) of all the samples without honey (Figure 3) can be considered similar (p > 0.05). Although the addition of natural polymers altered the degree of crystallinity of the samples, the amorphous chain entanglements and the crystallites formed in freeze-thawing maintained the samples’ structural integrity\cite{62}. The addition of honey decreased the gel fraction of the samples (p < 0.05). The gel fraction property is related to the hydrogel’s crosslinked polymers chains that remain insoluble when immersed in aqueous fluid. The PVA-H GF is significantly higher than the PVA-CMC-H GF which is, in turn, significantly higher than the PVA-G-H, GF (p < 0.05). The GF of the honey samples diminish in accordance with the decrease in the samples’ degree of crystallinity, indicating that the samples crystallites work as physical crosslinking points. The addition of the natural polymers, as well as the addition of honey, diminishes the gels’ degree of crystallinity, diminishing the gels’ GF\cite{63,64}.

The samples without honey presented similar weight loss (p > 0.05), Figure 3. The PV A samples weight loss (WL) could be due to biodegradation in saline: “hydrolytic cleavage of hydrogen bonding among -OH groups of PVA chains”, as discussed by Kamoun et al.\cite{65}.

### Table 2. Data regarding the samples degree of crystallinity ($X_c$), glass transition temperature ($T_s$), melting temperature ($T_m$), swelling degree (SD), gel fraction (GF), weight loss (WL), $S. aureus$ growth.

| Samples     | DSC ($T_s$) ($°C$) | $T_m$ (%) | $X_c$ (%) | SD | GF | WL | Growth (CFU/g) | Reduction (%) |
|-------------|--------------------|-----------|-----------|-----|----|----|----------------|---------------|
| PVA         | 87                 | 214       | 54        | 142 | 89 | 10 | 5,5 x 10$^6$   | 0             |
| PVA-H       | 80                 | 150       | 5         | 79  | 45 | 54 | 5,7 x 10$^6$   | 0             |
| PVA-CMC     | 69                 | 223       | 19        | 363 | 88 | 11 | 7,7 x 10$^6$   | 29.9          |
| PVA-CMC-H   | 85                 | -         | -         | 113 | 36 | 63 | 5,4 x 10$^6$   | 0             |
| PVA-G       | 79                 | 218       | 13        | 216 | 86 | 13 | 8,7 x 10$^6$   | 14.9          |
| PVA-G-H     | 102                | -         | -         | 49  | 22 | 77 | 7,4 x 10$^6$   | 0             |

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loss of the PVA blends could be related to PVA hydrolytic cleavage and also to the solubility of the natural polymers in aqueous media\(^75\). The samples containing honey presented significantly higher weight loss/biodegradation than the samples without honey. The WL followed the trend: PVA-H < PVA-CMC-H < PVA-G-H (\(p < 0.05\)), hypothesised to be due to degradation (due to chains leaching out by the saline solution) and honey delivery of both samples\(^{69}\). It is worth noting that only the three-layered samples were evaluated.

In this study, the PVA hydrogel can be considered a negative control (non-bactericidal) against gram-positive bacteria (\(S. \text{aureus}\)), allowing bacteria to grow in its presence\(^70\). The addition of honey inhibited the bacterial growth in the presence of the samples for PVA-CMC and PVA-G samples, Table 2. \(S. \text{aureus}\) attaches to proteins on the surface of collagen matrices and gelatin is a hydrolysed form of collagen\(^71\). Honey itself has a bactericidal effect due to its low water activity, the presence of \(\text{H}_2\text{O}_2\) and its low pH\(^72\). The antibacterial effect on \(S. \text{aureus}\) in wound healing hydrogels would depend on exceeding the minimum inhibitory concentration (MIC) of the honey with respect to the amount of honey in, and released from, the samples\(^73\). Several hydrogels containing honey present antibacterial activity\(^32,74\). The samples of the present work had 200 mg/mL of honey, but the MIC for this Brazilian honey might not have been reached for the PVA-honey samples. The PVA-CMC-H and PVA-G-H samples presented activity against \(S. \text{aureus}\) relative to their counterparts with no honey\(^59\). It was observed a bactericidal effect. It is known that even low concentrations of honey can stimulate wound healing\(^75\).

4. Conclusions

It was observed that the addition of some materials to PVA diminishes the gels crystallinity and gel fraction (related to the degree of crosslinking of the gels), altering the samples ability to swell, where the samples with honey presented lower fluid uptake than samples without it, since honey can occupy and obstruct pores. Nonetheless, the samples containing honey presented significantly higher biodegradation (hydrolytic degradation/weight loss) than the samples without honey. The samples with NaCMC or gelatin and honey were the ones that presented the highest activity against \(S. \text{aureus}\) relative to their honey-free counterparts, showing potential to be used as wound care materials.

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