Effects of grape seed extract on properties of type I collagen scaffolds

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Abstract: To obtain a material with potential for use in tissue engineering, anionic collagen was obtained from porcine serosa (S) and bovine tendon (T) by alkaline hydrolysis for 72h. Part of this collagen was mixed with water to obtain 4% (weight/weight) collagen suspension and part was solubilized in acetic acid pH 3.5 to obtain 1.5% (w/w) gel. The suspensions were mixed with their respective gels (2:1) (suspension: gel) and grape seed extract, whose main product is proanthocyanidin, was added at concentrations of 0.03% and 0.5%, thus obtaining the scaffolds SC (serosa collagen suspension and gel), TC (tendon collagen suspension and gel), SCP003 (SC with 0.03% extract), TCP003 (TC with 0.03% extract), SCP05 (SC with 0.5% extract added) and TCP05 (TC with 0.5% extract). The materials were analyzed by differential scanning calorimetry (DSC), scanning electron microscopy (SEM) and characterized by phosphorus buffered saline absorption assay and in vitro biological stability assay. By DSC it is observed that the addition of 0.5% of extract increases the denaturation temperature (Td) of collagen, indicating that at this concentration the extract acts as polymer crosslinking agent. SEM shows disorganized cross–section pores in all scaffolds, not exceeding 130 μm. Absorption and degradation assays indicated that the addition of 0.5% addition of 0.5% extract increases the absorption of phosphate buffered saline (PBS) by the scaffolds and decreases the degradation percentage by collagenase.

These results suggests that the scaffolds can be used for different applications, e.g. as hemostatic agent.

Keywords: Biopolymer; bovine tendon; porcine serosa; crosslinking.

Introduction

Topical absorbable hemostatic agents may be used to stop bleeding and retain wound exudate as they initiate platelet aggregation allowing blood coagulation. Different hemostatic agents have been developed and are available to assist surgeons in the treatment of bleeding. Among the various materials used are the microfibriillary collagen, bovine or porcine gelatin sponges, oxidized cellulose, gelatin sealant matrix, thrombin, etc.¹,²,³

Collagen and its derivatives are chemically attractive in wound healing, with good biological and hemostatic properties. In addition, they have biocompatible, low toxicity and biodegradable characteristics.¹ Its use as a hemostatic agent occurs due to its capacity for platelet aggregation and ability to initiate the cascade of blood coagulation.⁴,⁵

Collagen is the most abundant structural protein in the body of mammals. It is present in tendons, skin, blood vessels, intestines, bones and teeth and can be extracted from all these sources. The tendons are formed by dense regular connective tissue, which presents fibers arranged in a single direction giving tensile strength and low flexibility, unlike mature bone tissue, which presents collagen fibers arranged in the form of lamellae. Mucosa, muscularis, submucosa, and serosa layers form the intestine, which is composed of loose connective tissue, presenting collagen fibers without order.⁴,⁵

A chemical modification that can be performed with collagen is the selective hydrolysis of the carboxyamide groups of the asparagine and glutamine amino acid residues, leading to an increase of up to 134 negative charges per molecule.⁶ This modification allows the production of negatively charged collagen scaffolds at pH 7.4 with high biocompatibility and almost complete absence of chronic inflammatory response.¹,¹¹,¹²,¹³

Another modification that collagen may suffer is the crosslinking reaction, which increases the resistance to in vitro degradation and alters its mechanical properties.¹⁴ Proanthocyanidins are polyphenolic compounds in the category known as condensed tannins, which can be used as crosslinking agents due to the hydroxyl groups present in their structure, giving stability to hydrogen bonds and thus generating collagen structures with less biodegradability.¹⁵

Grape seed extract is a byproduct of the production of wines and sparkling wines, presenting several flavonoids such as anthocyanins, flavonols, catechins and proanthocyanidins, the most abundant. Proanthocyanidin has free radical scavenging ability, having superior antioxidant capacity than other oxidants like vitamin C, vitamin E, β-carotene. In addition, it has several pharmacological and biological properties, namely: cardioprotective, anti-tumor, bactericidal and anti-inflammatory.¹⁶

This study aims to obtain and characterize neutral scaffolds of anionic collagen obtained from two distinct sources (bovine tendon and porcine serosa), crosslinked with grape seed extract and having potential to be used as a hemostatic agent.

Materials and methods

Materials

Bovine tendon and porcine serosa were obtained at meat houses, and grape seed extract (Vitis sp.) was obtained from a drugstore.

Anionic collagen was obtained from two distinct sources, bovine tendon and porcine serosa, by the treatment in an aqueous alkaline solution (pH = 13) for 72 h using a protocol developed in previous studies in our laboratory.¹¹ In short, tendon or serosa were placed in an alkaline solution containing Ca(OH)₂, KOH, NaOH, Na₂SO₄, CaSO₄, KCl and NaCl for 72 h. Subsequently, the material was equilibrated for 6 h in a solution containing the same salts. Excess salts were removed by rinsing in 3% (w/w) boric acid, deionized water, 0.3% (w/w) EDTA solution, at pH 11 and deionized water up to pH 7.

Treated porcine serosa in water was ground in a blender and later its concentration was defined by lyophilization. Subsequently, it was divided in two parts, the first one was used to prepare a 4% suspension in water and the second one was solubilized in acetic acid pH 3.5 to obtain 1% collagen gel.

Treated bovine tendon was lyophilized and grounded in a blender. It was later divided in two parts, the first of which was suspended in water to obtain a 4% suspension and the other one was solubilized in acetic acid pH 3.5 to obtain 1% collagen gel.

The scaffolds were obtained by mixing the suspension with the collagen gel from the same source, in a 2:1 ratio (suspension: gel) and adding

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or not grape seed extract, which was previously solubilized in 20 mL of 50 % hydroethanolic.

Concentrations of 0.03 % and 0.5 % of grape seed extract were chosen based on previous studies in our laboratory.

The mixtures were placed in Teflon® molds and then lyophilized. They were neutralized using ammonia vapor for a period of 24 h and subsequently left under constant air flow to remove excess ammonia for 72 h. Scaffolds were nominated as shown in Table 1.

<table>
<thead>
<tr>
<th>Scaffold</th>
<th>Composition</th>
</tr>
</thead>
<tbody>
<tr>
<td>SC</td>
<td>serosa suspension and its gel</td>
</tr>
<tr>
<td>SCP03</td>
<td>serosa suspension and its gel + grape seed extract 0.03% (w/w)</td>
</tr>
<tr>
<td>SCP05</td>
<td>serosa suspension and its gel + grape seed extract 0.5 % (w/w)</td>
</tr>
<tr>
<td>TC</td>
<td>tendon suspension and its gel</td>
</tr>
<tr>
<td>TCP03</td>
<td>tendon suspension and its gel + grape seed extract 0.03% (w/w)</td>
</tr>
<tr>
<td>TCP05</td>
<td>tendon suspension and its gel + grape seed extract 0.5 % (w/w)</td>
</tr>
</tbody>
</table>

Table 1 — Scaffolds denomination

Methods

Differential scanning calorimetry (DSC)

DSC curves were obtained with a computer–interfaced differential scanning calorimeter (Model DSC 2010, TA Instruments, New Castle, DE, U.S.A.). Samples of 20 mg were sealed in hermetic aluminum pans and equilibrated at the initial temperature. The scanning was carried out with a heating rate of 10°C min-1 from 5 to 120 °C and performed under a nitrogen atmosphere. Denaturation temperature (Td) was determined as the inflection point value of the corresponding endothermic effect.

Scanning Electron Microscopy (SEM)

Samples were coated with a thin layer of gold of 6 nm. The specimens were examined with a Leo 440, LEO Electron Microscopy Ltd. (Cambridge, England) with an accelerating voltage of 20 kV. For the cross section analysis, the scaffolds were frozen in liquid nitrogen, fractured and lyophilized. To measure the pore size, ImageJ software was used and for each scaffold, 31 determinations were made using the approximation of the Martin's diameter and an image magnification of 500×.18

Absorption kinetics in phosphate buffered saline (PBS)

Scaffolds were placed in vials containing phosphate buffered saline (PBS). At specific time intervals, scaffolds were removed from the PBS and the excess of solution was removed using a 2 cm x 2 cm filter paper. The scaffolds were weighed and returned to the vials for further time taken. The process was made in quintuplicate. The amount of PBS absorbed was calculated by averaging the results found using equation (1), where \( w_{\text{swm}} \) is the mass of the swollen scaffold and \( w_{\text{dry}} \) is the mass of the scaffold before swelling.

\[
\% \text{ absorption} = \frac{(w_{\text{swm}} - w_{\text{dry}})}{w_{\text{dry}}} \times 100
\]  

In vitro biological stability (collagenase)

For in vitro biological stability determination, collagenase solution in 10 mmol L-1 tris–HCl buffer at pH 7.4 containing 25 mmol L-1 CaCl2 was prepared. Collagenase solution (10 U mg-1 of collagen) in tris–HCl buffer, was added to scaffolds known weights which were subsequently placed in a bacteriological oven at 37 °C for 2 h. After this, the samples were washed deionized water, frozen and lyophilized to constant weight. The percentage of degraded collagen (% degradation) was determined by the difference in collagen mass before (\( w_{\text{initial}} \)) and after enzymatic degradation (\( w_{\text{final}} \)) by equation (2), and at least three determinations were made for each scaffold.

\[
\% \text{ degradation} = \frac{(w_{\text{initial}} - w_{\text{final}})}{w_{\text{initial}}} \times 100
\]

Statistical Analysis

The analysis results were analyzed using ANOVA. A value of p<0.05 was considered statistically significant.

Results and discussion

The presence of grape seed extract is noted for its beige color in TCP003 and SCP003 scaffolds and brown in TCP05 and SCP05 scaffolds. The scaffolds without extract (TC and SC) are white and all of them presented touch–resistant appearance and reversibility to small deformations.

DSC curves showed a transition that appears as a discontinuity at baseline, giving the collagen denaturation temperature (Td). Denaturation occurs due to heating and can be defined as the transition of the collagen triple helix to a form in which intramolecular bonds are broken down, changing from a highly organized structure to a disorganized state called gelatin.19 The values of collagen denaturation temperatures are shown in Table 2.

Native tissues (tendon and serosa) had a denaturation temperature of 63.1°C, values close to those found by Willett20 and Hizaji21, who analyzed the Td of bovine tail tendon collagen and obtained values of 62.4°C and 63.3°C, respectively. Hirata22 evaluated the Td of bovine serum and obtained a value of 64.6°C, close to those found in this study.

Therefore, a reduction in Td of 9.8 and 3.2°C is observed for native tissues compared to their respective TC and SC scaffolds, respectively. This indicates that alkaline treatment destabilizes the triple collagen helix, and that this destabilizing effect is greatest in tendon matrices.

Alkaline hydrolysis also modifies the macromolecular aggregation of native collagen23, which is different for tendon and serosa tissues. Differences in Td of SC and TC may indicate that the structure of the serosa is more resistant to destabilization, keeping part of its intrinsically cross–linked loose connective tissue structure.

The denaturation temperature of TC and TCP03 are very close, as well as for SC e SCP003 suggesting that crosslinking by grape seed extract was not effective at 0.03 % concentration.

Scaffolds containing 0.5 % of extract had an increase in denaturation temperature when compared to the scaffolds without extract, being of 14.7°C for the bovine tendon scaffolds and 6.4°C for the serosa scaffolds, indicating that in both cases 0.5 % of extract is effective as a crosslinking agent.

In addition, the 0.5% extract cross–links collagen to the extent that the Td of SCP05 and TCP05 is larger than that of their respective originating tissues. This is because there are still crosslink available sites in the tendon and serosa and the proanthocyanidin present in the extract binds to available sites of the aminoacid proline group present in collagen, causing crosslinking.24

Scanning electronic microscopy allows to define the morphology of the scaffolds microstructures, such as pore size, distribution, porosity and interconnectivity, which are major factors in the interaction of the scaffolds with physiological environment, promoting or not the absorption of physiological liquids.
Photomicrographs (Figures 1–6) show the presence of surface pores and in the cross-section for all samples. Table 3 shows the average pore size, with the respective standard deviations (sd) and the coefficient of variation (Cv).

No statistical difference was observed between TC and TCP05 pore sizes, indicating that the extract addition is not a major factor in pore size in bovine tendon scaffolds. However, for serosa scaffolds, the addition of extract at concentrations of 0.03 % and 0.5 % causes the emergence of pores on the scaffolds surface.

The histograms show that TC scaffold (Figure 2), with 74.0% Cv, has a bimodal system with a pore size frequency between 10–20 μm and 30–40 μm. In the case of the SCP003 scaffold (Figure 3), it also has a bimodal system with higher pore size frequencies between 20–30 μm and 30–40 μm. The other scaffolds have systems with one mode, and for the scaffolds TCP003 (Figure 4), SCP05 (Figure 5) and TCP05 (Figure 6) the largest frequency of pore size occurs between 40–50 μm, 20–30 μm, and 30–40 μm, respectively. In the cross section, it is observed that only the TCP003 scaffold has a bimodal system, with a higher incidence of pores between 40–50 μm and 50–60 μm. Only two scaffolds showed statistical difference between the pore size of the surface and the cross section. SC scaffold showed no pores on the surface and SCP05, which presented great pore variation in its cross section, presenting Cv of 54.2%.

Nyberg (2017) and Zhang (2018) show that pores smaller than 100 μm facilitate the diffusion of physiological fluids within the scaffold and promote a capillary force that improves cell attachment in the periphery of the scaffold and can induce cell penetration.

The ability of a scaffold to retain water and electrolytes is an important property for evaluating its effectiveness as a hemostatic material, so the absorption test is essential. Absorption of saline phosphate buffer by collagen scaffolds arises due to various electrostatic interactions. When the first water molecules enter the pores of the scaffold, they bind (hydrate) the hydrophilic groups present in collagen and grape seed extract.

**Table 2 – Denaturation temperature (Td).**

<table>
<thead>
<tr>
<th>Scaffold</th>
<th>Td (°C)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Bovine tendon</td>
<td>63.1</td>
</tr>
<tr>
<td>Porcine serosa</td>
<td>63.1</td>
</tr>
<tr>
<td>TC</td>
<td>53.3</td>
</tr>
<tr>
<td>SC</td>
<td>59.9</td>
</tr>
<tr>
<td>TCP003</td>
<td>53.7</td>
</tr>
<tr>
<td>SCP003</td>
<td>60.8</td>
</tr>
<tr>
<td>TCP05</td>
<td>68.0</td>
</tr>
<tr>
<td>SCP05</td>
<td>66.3</td>
</tr>
</tbody>
</table>

**Figure 1 – SC scaffold: (A) surface pore size distribution histogram; (B) surface photomicrograph; (C) cross-sectional pore size histogram (D) cross section photomicrograph.**
Figure 2 – TC scaffold: (A) surface pore size distribution histogram; (B) surface photomicrograph; (C) cross-sectional pore size histogram (D) cross section photomicrograph.

Figure 3 – SCP003 scaffold: (A) surface pore size histogram; (B) surface photomicrograph; (C) cross-sectional pore size histogram (D) cross section photomicrograph.
Figure 4 – TCP003 scaffold: (A) surface pore size distribution histogram; (B) surface photomicrograph; (C) cross-sectional pore size histogram (D) cross section photomicrograph.

Figure 5 – SCP05 scaffold: (A) surface pore size histogram; (B) surface photomicrograph; (C) cross-sectional pore size histogram (D) cross section photomicrograph.
adhesive interactions between the various polar aminoacids residues present in collagen and PBS are stronger than the cohesive forces of the buffer itself. This causes the buffer to be absorbed throughout the scaffold, causing it to swell. PBS absorption at 30, 60 and 1200 min is shown at Table 4.

Figure 7 shows the percentage of absorption at different times. TCP05 and TCP003 had their maximum absorption at 2750 % in the first minute, different of TC, which had maximum absorption at 1920% after 1200 minutes.

Serosa scaffolds absorbed their maximum around 30 minutes, which SC and SCP003 had statistically equal absorption around 1400% and the SCP05 absorbed around 2750 %.

Scaffolds TCP05 and SCP05 have statistically equal absorption around 2700%, almost 2 times higher than the scaffolds without extract. This is due to the large number of hydroxyl groups present in proanthocyanidin, which make the material most hydrophilic.

Swelling of a material is hampered by inter and intramolecular bonding, intrinsic stiffness and degree of crosslinking. Then, a difference in the absorption of the TC and SC was expected, since the serosa scaffold showed higher thermal stability, which may be due to a larger number of bonds in its structure (intrinsic cross-linking).

This difference is only observed between TCP003 and SCP003, since the presence of 0.03 % extract is not able to crosslink collagen to the point of increasing its Td, but it contributes to the hydrophilicity of the material, making the scaffold TCP003 more hydrophilic than SCP003, which is difficult to absorb due to its intrinsic cross-linking.

### Table 3 – Surface and cross section pore size of the scaffolds.

<table>
<thead>
<tr>
<th>Scaffolds</th>
<th>Surface</th>
<th>Cv (%)</th>
<th>Cross section</th>
<th>Cv (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>TC</td>
<td>41.1 ± 30.4 a,b</td>
<td>74.0</td>
<td>43.2 ± 13.7 b,d</td>
<td>30.3</td>
</tr>
<tr>
<td>SC</td>
<td>31.1 ± 17.8 b</td>
<td>57.1</td>
<td>48.4 ± 26.2 b,c</td>
<td>54.2</td>
</tr>
<tr>
<td>TCP003</td>
<td>52.9 ± 21.0 a</td>
<td>39.7</td>
<td>56.7 ± 22.2 a,b</td>
<td>39.2</td>
</tr>
<tr>
<td>SCP003</td>
<td>36.4 ± 16.6 b</td>
<td>45.5</td>
<td>38.3 ± 9.3 c,d</td>
<td>24.2</td>
</tr>
<tr>
<td>TCP05</td>
<td>41.9 ±19.5 a,b</td>
<td>46.4</td>
<td>41.2 ±17.5 c,d</td>
<td>42.5</td>
</tr>
<tr>
<td>SCP05</td>
<td>32.4 ± 14.5 b</td>
<td>44.6</td>
<td>63.1 ± 23.8 a</td>
<td>37.7</td>
</tr>
</tbody>
</table>

In the same column, same letter means statistically equal numbers (p<0.05) sd: standard deviations. Cv: coefficient of variation (%).
linking as observed by their denaturation temperature, only TC, SC, TCP05 and SCP05 were used in the degradation assays, shown in Table 5.

The scaffold TC presented the higher percentage of degradation, being 74.0 ± 5.0 and, therefore, having lower biological stability in vitro. The scaffold SC presented a degradation percentage of 9.3 ± 2.1, indicating greater stability of serosa collagen against collagenase, probably due to the higher number of intrinsic serosa tissue crosslinks, as already mentioned.

With the addition of 0.5% extract, the percentage of tendon scaffolds degradation decreased from 74.0 ± 5.0 to 15.3 ± 0.4 due to crosslinking caused by proanthocyanidin and other flavonoids present in grape seed extract.

Degradation percentage for SC and SCP05 were expected to be statistically different, however this was not observed. Maybe another test using a higher degradation time could result in a difference of degradation between serosa scaffolds.

Ma et al. evaluated the biodegradability of collagen of 0.25% glutaraldehyde crosslinked bovine tendon observing that there was a reduction of biodegradability of approximately 87 %, close to the observed in this study with tendon scaffold that when cross–linked with 0.5 % extract showed reduction of approximately 80 %.

Rodrigues also evaluated trypsin enzymatic stability of glutaraldehyde cross-linked collagen. Glutaraldehyde cross–linking reduces the degradation percentage of collagen scaffolds too. This indicates a possible substitution of glutaraldehyde for the extract, as glutaraldehyde presents high cytotoxicity.

<table>
<thead>
<tr>
<th>Scaffold</th>
<th>Absorption (%) ± sd</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>30 min</td>
</tr>
<tr>
<td>TC</td>
<td>1575 ± 217 b</td>
</tr>
<tr>
<td>SC</td>
<td>1374 ± 131 b</td>
</tr>
<tr>
<td>TCP03</td>
<td>2739 ± 192 a</td>
</tr>
<tr>
<td>SCP03</td>
<td>1469 ± 106 b</td>
</tr>
<tr>
<td>TCP05</td>
<td>2342 ± 305 a</td>
</tr>
<tr>
<td>SCP05</td>
<td>2813 ± 580 a</td>
</tr>
</tbody>
</table>

In the same column, same letter means statistically equal numbers (p<0.05) sd: standard deviations.

Table 4 – PBS absorption kinetics (%).

<table>
<thead>
<tr>
<th>Scaffold</th>
<th>Degradation (%) ± sd</th>
</tr>
</thead>
<tbody>
<tr>
<td>TC</td>
<td>74.0 ± 5.0 a</td>
</tr>
<tr>
<td>SC</td>
<td>9.3 ± 2.1 b, c</td>
</tr>
<tr>
<td>TCP05</td>
<td>15.3 ± 0.4 b</td>
</tr>
<tr>
<td>SCP05</td>
<td>3.7 ± 2.5 c</td>
</tr>
</tbody>
</table>

In the table, same letter means statistically equal numbers (p<0.05) sd: standard deviations.

Table 5 – Scaffolds degradation ± sd.
Conclusion
Photomicrographs showed the presence of surface pores and in the cross-section for all samples, not larger than 130 μm. Cross-linking by 0.5% extract increases collagen denaturation temperature, reduces degradation percentage 12 and increases the swelling, which may result in scaffolds for different applications such as hemostatic agent or soft tissue regeneration.

Conflict Of Interests
There are no conflicts to declare.

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References


