Histological Evaluation of Direct Pulp Capping with Silk Sericin: A Preliminary Animal Study

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Introduction: This histological study analyzed silk sericin as a potential direct pulp capping biomaterial in contact with pulp and comparing its response to calcium hydroxide. Methods and Materials: Twenty maxillary first molars from Wistar male rats were used, with 60 days of age, between 200 and 300 gr, which were divided in 4 groups (n=5): G1 and G3, controls, capped with calcium hydroxide in 7 and 30 days, respectively; G2 and G4, capped with silk sericin in 7 and 30 days, respectively. Circular cavities were prepared for pulp exposure, where capping materials were applied, being posteriorly restored with glass ionomer cement. After completion of each observation period, the animals were sacrificed and molars were histologically processed for analysis in light microscopy to evaluate presence of necrosis in pulp tissue, inflammatory cells infiltration and tertiary dentin formation. Data analysis was carried out using Kruskal-Wallis and Dunn’s post hoc tests.

Results: After 7 days, there was less necrosis and inflammatory cells infiltration in G1 when compared to G2 (P=0.007 and P=0.008, respectively). After 30 days, a sample of G3 induced tertiary dentin formation and G4 showed decrease in inflammation (P=0.041) compared to G2.

Conclusion: Among the determined experiment conditions, it was concluded that contact between silk sericin and pulp tissue showed improved inflammatory response throughout treatment and new cells proliferation. However, silk sericin adhesion in pure form did not show capability for induction of tertiary dentin formation.

Keywords: Biocompatible Materials; Calcium Hydroxide; Dental Pulp Capping; Sericin

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The sericin powder on a glass plate and Angelus MTA applicator of 0.6 mm in diameter can be seen.

Figure 1.

The operating table for immobilization and conservation of the animal’s mouth opening.

Figure 2.

Regarding calcium hydroxide, its effect on pulp capping is seen as the result of chemical damage caused by hydroxyl ions liberation. Initially, calcium hydroxide application to exposed pulp causes a development of superficial necrosis and mild inflammation due to its alkaline pH, stimulating pulp defense and tissue healing to form a restorative dentin bridge through cell differentiation, extracellular matrix secretion and subsequent mineralization [18].

Based on these considerations, it is opportune to analyze silk sericin as a potential biomaterial in contact with dental pulp, histologically comparing its response to direct pulp capping in mice with calcium hydroxide, so that it is possible to evaluate the presence of necrosis on pulp tissue, infiltration of inflammatory cells and tertiary dentin formation during two periods of time: 7 and 30 days.

Materials and Methods

The experimental procedures of this paper are in accordance with the Ethical Principles on Animal Experimentation, employed by the Brazilian College of Animal Experimentation (COBEA) and were analyzed and approved by the Ethics Committee on Animal Use (CEUA) of Universidade Estadual do Oeste do Paraná under protocol # 60/17.

This study was performed on 20 maxillary first molars of 60-day male mice (Rattus norvegicus albinus, Wistar), weighing about 200 to 300 gr, held in a controlled ambient, bright-dark cycle of 12 h, room temperature of 21±1°C, water and pasty ration ad libitum. These molar teeth were histologically evaluated under two periods of time: 7 and 30 days distributed equally and randomly in 4 groups (n=5), being group 1 (control): calcium hydroxide in 7 days; group 2: silk sericin in 7 days; group 3 (control): calcium hydroxide in 30 days; group 4: silk sericin in 30 days.
Silk sericin preparation
Silk sericin protein was extracted from silkworm cocoons (Bombyx mori), obtained from the sericulture company (BRATAc Silk do Brasil®, Londrina, Paraná, Brazil). Cocoons were cut into small pieces (about 1 cm²) and the equivalent of 6 gr of cocoons were submerged in 100 mL of distilled water, in a 500 mL Erlenmeyer. The solution was autoclaved, (autoclave CS 30–Prismatec) at 120°C and pressure of 1 kgf/cm², for 60 min. The raw extract was then filtered with an 18 mesh sieve, to separate/ remove fibroin, from which a silk sericin hydrolyzed was obtained. This protocol was based on studies by Gimenes et al. [19], who utilized water, without any chemical additive, for silk sericin extraction. The hydrolyzed silk sericin was then frozen and freeze-dried (Liofilizador LT 1000, Terroni Equipamentos Ltda®, São Carlos, SP, Brazil), for approximately 54 h, obtaining silk sericin powder, stored in room temperature until its utilization (Figure 1).

Direct pulp capping
Animals were weighted previously and anesthetized with a combination of ketamine 10% 0.1 mL/100 gr of weight (Dopalen injétável®, Ceva Saúde Animal, Paulínia, SP, Brazil) and 2% xylazine 0.05 mL/100 gr of weight (Anasedan injétável®, Ceva Saúde Animal, Paulínia, SP, Brazil), through an intraperitoneal injection, and were positioned on appropriate operating table [20], which allows wide mouth opening, enabling access to teeth in maxillary region (Figure 2). Circular cavities were prepared on the maxillary first molars, at the occlusal surface above mesial central surface of the teeth, aided by an endodontic motor VDW Silver (VDW®, Munich, Germany) at 400 rpm, with a ¼ round diamond bur (KG Sorensen Ind & Com®, Alphaville, São Paulo, Brazil), which was used under microscopic magnification and saline solution watering until bleeding was obtained. The diameter of exposed pulp area ranged from 0.5 to 0.7 mm. Pulp hemorrhage is controlled and cavity is dried with sterilized paper cones, adapted from Shinkai et al. [21]. Materials were applied using Angelus MTA applicator of 0.6 mm in diameter metallic device (Angelus®, Londrina, PR, Brazil) and cavities were sealed with Ketac Molar Easymix glass ionomer cement (3M ESPE®, São Paulo-SP, Brazil) (Figure 3).

Samples obtaining
After completion of each observation period (7 and 30 days after pulp capping), the animals were sacrificed, using intraperitoneal injections with an overdose of the same anesthetics used in operating procedure (four times greater). Maxilla was then removed and properly labeled, stored in glass container and underwent fixation in 10% formaldehyde solution for 48 h, and later in 70% alcohol for 15 days.

Histological processing
Heminaxilla was decalcified with 5% trichloroacetic acid (TCA), in room temperature for 20 days. Pieces were analyzed to evaluate expected decalcification degree, with renewal of the
TCA solution every 5 days. After decalcification, pieces were rinsed under running tap water and underwent automated histological processing (Automated tissue processor, Leica Microsystems TP1020®, Nussloch, Germany). After that, pieces were embedded in paraffin and blocks were obtained (Purified Paraffin, Vetec Química Fina®, RJ, Brazil). These blocks were cut with semi-automatic microtome (Hestion®, ERM3000, Daintree Scientific, St. Helens, Australia) in order to obtain sections of 5 µm in width. Cuts were selected according to region of exposure and mounted in histological lamina with hematoxylin and eosin stain technique.

**Histological analysis**

Histological analyses were performed by two previously trained observers, in blind fashion, using 40 ×, 100× and 400× magnifications (Optic Microscope Leica Microsystems® ICC50HD, Nussloch, Germany) and image capture system (Las Ez–Leica®, version 2.10, 2012, Solms, Germany). Descriptive analysis was performed, analyzing the following parameters: presence of pulp necrosis, infiltration of inflammatory cells and tertiary dentin formation, which were classified using scores and are described in Table 1, adapted from Suzuki et al. [22].

**Statistical analysis**

For the evaluation of variables of this study, all scores were expressed as median±interquartile range. Histological comparisons and sample calculation were based on study by Shinkai et al. [21] and performed with BioStat 5.3 program (Mamirauá Institute, Belém, Pará, Brazil), using Kruskal-Wallis test and Dunn’s post hoc test, with significance level of 0.05. Furthermore, as few animals as possible were used per group, for animal protection.

**Table 1.** Scores and parameters analyzed

<table>
<thead>
<tr>
<th>Scores</th>
<th>Parameter</th>
</tr>
</thead>
<tbody>
<tr>
<td>0</td>
<td>Absence of necrosis</td>
</tr>
<tr>
<td>1</td>
<td>Presence of necrosis on coronary third of the pulp</td>
</tr>
<tr>
<td>2</td>
<td>Presence of necrosis on coronary third and middle</td>
</tr>
<tr>
<td>3</td>
<td>Presence of necrosis on all the pulp</td>
</tr>
<tr>
<td>0</td>
<td>Absence or occasional presence of inflammatory cells on pulp (none)</td>
</tr>
<tr>
<td>1</td>
<td>Mild/chronic inflammation (mild)</td>
</tr>
<tr>
<td>2</td>
<td>Inflammatory reactions such as micro abscesses or polymorphonuclear leukocytes infiltrates, histiocytes and lymphocytes affecting the coronary pulp up to half of the root pulp (moderate)</td>
</tr>
<tr>
<td>3</td>
<td>Inflammatory reaction affecting all of the pulp tissue (severe)</td>
</tr>
<tr>
<td>4</td>
<td>Unable to evaluate inflammatory cells due to necrotic degeneration</td>
</tr>
<tr>
<td>0</td>
<td>Complete formation of the dentinal bridge (complete)</td>
</tr>
<tr>
<td>1</td>
<td>Partial/incomplete dentinal bridge formation, which extends to more than half of the exposure region, but doesn’t completely close the exposure region (partial)</td>
</tr>
<tr>
<td>2</td>
<td>Initial dentinal bridge formation, extending to no more than half of the exposure region (initial)</td>
</tr>
<tr>
<td>3</td>
<td>Without dentinal bridge formation (none)</td>
</tr>
</tbody>
</table>

**Table 2.** Statistical analysis of presence of necrosis on pulp tissue for the evaluated materials (median ± interquartile range).

<table>
<thead>
<tr>
<th>Material/Time</th>
<th>7 days</th>
<th>Scores</th>
<th>Frequency</th>
<th>30 days</th>
<th>Scores</th>
<th>Frequency</th>
</tr>
</thead>
<tbody>
<tr>
<td>Calcium Hydroxide</td>
<td>1 ± 0.0 b b</td>
<td>1</td>
<td>4</td>
<td>3 ± 1.0 b b</td>
<td>3</td>
<td>1</td>
</tr>
<tr>
<td></td>
<td>2</td>
<td>1</td>
<td>1</td>
<td>3 ± 1.0 b b</td>
<td>1</td>
<td>3</td>
</tr>
<tr>
<td></td>
<td>3</td>
<td>0</td>
<td>0</td>
<td></td>
<td></td>
<td>3</td>
</tr>
<tr>
<td>Sericin</td>
<td>3 ± 0.0 b b</td>
<td>1</td>
<td>0</td>
<td>2 ± 1.0 b a</td>
<td>2</td>
<td>1</td>
</tr>
<tr>
<td></td>
<td>2</td>
<td>1</td>
<td>1</td>
<td>3 ± 1.0 b b</td>
<td>3</td>
<td>3</td>
</tr>
<tr>
<td></td>
<td>3</td>
<td>4</td>
<td>3</td>
<td></td>
<td></td>
<td>2</td>
</tr>
</tbody>
</table>

Different upper case letters in the same column mean statistically significant differences, P<0.05. Different lower case letters in the same line mean statistically significant differences, P<0.05.
Results

Seventh day evaluation
In response to calcium hydroxide on the seventh day (group 1), it was observed that the superficial portion of the odontoblastic layer and coronary pulp adjacent cells were disorganized due to the diamond bur action during pulp exposure. Furthermore, necrosis was observed on pulp tissue at cervical third of the pulp in 80% of the samples, and acute-to-moderate inflammatory infiltrate involving coronary pulp up to half of the root pulp, represented by neutrophil exudation areas. On the deep layers pulp tissue had moderate cellularity with fibroblasts, granulation tissue and hyperemia in the majority of analyzed samples (Figure 4A).

On the other hand, pulp response to silk sericin in 7 days (group 2) displayed necrotic degeneration involving all coronary and root pulp in 80% of the cases, rendering qualitative morphological analysis of other parameters impossible (Figure 4B). It can be noted that, in Tables 2 and 3 respectively, group 1 had better score comparing to group 2, regarding pulp tissue necrosis degree ($P=0.007$) and presence of inflammatory cells infiltration ($P=0.008$). Concerning tertiary dentin formation at 7 days, there was no dentin barrier formation in all cases, regardless of material used, with no statistical difference between groups for this criterion ($P>0.05$).

Thirtieth day evaluation
Within 30 days, presence of necrosis was observed predominantly on coronary, middle and apical thirds of pulp tissue for the calcium hydroxide group (group 3). Regarding silk sericin (group 4), presence of necrosis was observed mainly on coronary third and middle of pulp. Underneath necrosis, for both groups, chronic moderate inflammatory infiltrate was observed and, in some cases, extensive, characterized mainly by mononuclear leukocytes, such as lymphocytes and macrophages. Regarding tertiary dentin formation, in only one sample from group 3 it was possible to see partial atubular dentin barrier formation capability, in more than half of the exposure region (Figure 4C), while group 4 did not show any tertiary dentin barrier formation (Figure 4D). In comparison between groups, there was no significant statistical difference for any of the three evaluation criteria ($P>0.05$).

Comparison between 7 and 30 days
Regarding the occurrence of pulp necrosis on groups that underwent calcium hydroxide treatment (group 1 and group 3), it was possible to note that time duration was a spread aggravating factor ($P=0.010$), since pulp area affected by necrosis was greater in 30 days (Table 2). Concerning inflammatory cells infiltration in the same groups, there was also a greater degree of inflammation ($P=0.012$) after 30 days (Table 3). Concerning tertiary dentin formation, calcium hydroxide induced dentin barrier formation in 20% of samples in 30 days. It was not observed statistically significant differences among the groups regarding the different time periods (Table 4). Silk sericin had decreased levels of inflammation after 30 days of treatment ($P=0.041$), even though it did not show histological signs of mineralized material formation. Silk sericin did not show significant statistical difference between 7 and 30 days concerning pulp tissue necrosis and tertiary dentin formation ($P>0.05$).

Table 3. Statistical analysis of inflammatory cells infiltration for the evaluated materials (median±interquartile range)

<table>
<thead>
<tr>
<th>Material/Time</th>
<th>7 days Scores</th>
<th>Frequency</th>
<th>30 days Scores</th>
<th>Frequency</th>
</tr>
</thead>
<tbody>
<tr>
<td>Calcium hydroxide</td>
<td>2±0.0 Ab</td>
<td>1</td>
<td>4±1.0 Ab</td>
<td>1</td>
</tr>
<tr>
<td>Sericin</td>
<td>4±0.0 Bb</td>
<td>1</td>
<td>2±1.0 Ab</td>
<td>1</td>
</tr>
</tbody>
</table>

Different upper case letters in the same column mean statistically significant differences, $P<0.05$. Different lower case letters in the same line mean statistically significant differences, $P<0.05$

Table 4. Statistical analysis of tertiary dentin formation for the evaluated materials (median±interquartile range)

<table>
<thead>
<tr>
<th>Material/Time</th>
<th>7 days Scores</th>
<th>Frequency</th>
<th>30 days Scores</th>
<th>Frequency</th>
</tr>
</thead>
<tbody>
<tr>
<td>Calcium hydroxide</td>
<td>3±0.0 Ab</td>
<td>1</td>
<td>3±0.0 Ab</td>
<td>1</td>
</tr>
<tr>
<td>Sericin</td>
<td>3±0.0 Ab</td>
<td>1</td>
<td>3±0.0 Ab</td>
<td>1</td>
</tr>
</tbody>
</table>

Different upper case letters in the same column mean statistically significant differences, $P<0.05$. Different lower case letters in the same line mean statistically significant differences, $P<0.05$
Discussion

In this study, pulp exposure treatment with lyophilized silk sericin in pure form pointed its pro-inflammatory capacity during initial response and has also shown to be more aggressive to pulp tissue in 7 days of treatment when compared to calcium hydroxide. On the other hand, inflammatory response decreased after 30 days, fact that can be compared to a study by Aramwit et al. [13] which concluded that healing process of wounds showed improvement in treatment after use of silk sericin, demonstrating that inflammatory mediators TNF-α and IL-1β, when the experiment performed with silk sericin, were significantly lower.

Expectations on potential use of silk sericin as biomaterial in odontology, as a complement or even as a new alternative in direct pulp capping, is based on observation of results on this biomaterial application in other tissues, where it presented itself to be very promising [6-10, 13]. Considering that there are no studies using this protein for such end, these are the first morphological and functional data obtained from silk sericin application directly on exposed pulp.

The initial aggressive response may be due to the fact that glass ionomer cement has acid and caustic properties [23] and may have caused injury to pulp, leading to an exacerbated inflammatory response.

For many years, importance of inflammation in pulp healing was underestimated, being considered only as an undesirable effect. However, there are evidences that inflammation is a requirement for tissue healing and pulp regeneration, and immuno-competent cells (monocytes, macrophages and stem cells) migrate towards the crown as part of this process. Due to the capping agent’s high alkalinity, mineralization starts and becomes thicker, while inflammatory processes induce pulp cells proliferation, presenting cells both greater in number and size [24].

Corroborating study by Aramwit et al. [6] and Ersel et al. [9], showed that silk sericin accelerated skin tissue regeneration and supported new cells proliferation, its contact with dential pulp. In this study, while not capable of inducing tertiary dentin formation and not suggesting interaction with odontoblasts, showed a lower degree of inflammatory cells infiltration after 30 days of treatment when compared to its initial response, demonstrating improvement during treatment and indicating a reparative response tendency. Since it did not form tertiary dentin, lyophilized silk sericin applied in pure form did not show direct interaction with odontoblasts, although further studies could test its reparative potential when associated to materials that induce hard mineralized tissue formation, contributing to inflammatory response improvement.

Despite its potential, pure silk sericin forms fragile filaments and it becomes difficult to use it as a biomaterial in tissue engineering [25]. Thus, different strategies have been applied in order to increase its physical properties [7]. In fact, improving some properties of the material may have deleterious effects on other properties. Therefore, clinical effectiveness of the modification in a given material in addition to its physical properties, antibacterial activity, sealing ability and biocompatibility should be considered [26]. Additionally, cavity size performed on animal tooth, as well as pulp exposure extension, hinder material insertion and settlement efficiency on desired region.

Calcium hydroxide is a material that holds a long history in clinical use, considered the gold standard for this application, for many years, and still is widely utilized for exposed pulp direct coverage [4, 22]. In this study, direct pulp capping performed with calcium hydroxide showed considerable increase in necrosis incidence and pulp inflammation after 30 days of treatment when compared to its initial response, in 7 days, in addition to showing tertiary dentin formation capacity. This result revealed to be similar to a study reported by Tran et al. [27], who observed a persistent necrotic layer in all groups tested with calcium hydroxide next to exposure region, and capacity for a barrier formation of mineralized dentin under the capping material, often incomplete and predominantly porous.

Tertiary dentin formation in relation to pulp capping with calcium hydroxide after 30 days was noted in only 20% of samples. This percentage could not be more significant due to reduced number of samples, as it is a well-established material in dental practice. Nonetheless, this result displays repair capability under calcium hydroxide activity, as already noted in several similar studies [18, 22, 23, 28].

Conclusion

Based on the results, it was possible to conclude that silk sericin contact with pulp tissue showed improvement in inflammatory response throughout treatment and new cells proliferation. However, silk sericin adhibition in pure form did not show capability for induction of tertiary dentin formation. When compared to calcium hydroxide, initial response to silk sericin displayed higher degree of necrosis and inflammatory cells infiltration, decreasing after 30 days. Therefore, considering all silk sericin properties and benefits extensively related on literature, more researches should be performed, combining it with other similar biomaterials, or currently commercial materials for pulp capping.
Acknowledgment

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Conflict of Interest: 'None declared'.

References