Indonesian propolis suppressed the expression of COX-2 in inflamed rat dental pulp in direct capping treatment

Ardo Sabir*

Abstract

Objective: The aim of the present study was to analyze the expression of COX-2 on inflamed rat dental pulp after capping with propolis. Material and Methods: Propolis was obtained from South Sulawesi, Indonesia. Flavonoid and non-flavonoid were purified from an ethanolic extract of propolis (EEP). Eighty male Sprague Dawley rats were divided randomly into 5 groups, each group consisting of 16 rats. Group I as a negative control, rats were left without any treatment. A Class I cavity was prepared on the occlusal surface of permanent maxillary right first molar. Dental pulp was exposed and allowed in oral environment for 1 hour, then capping with EEP (group II), flavonoid propolis (group III), non-flavonoid propolis (group IV), or calcium hydroxide (Ca(OH)2) (group V). All cavities were filled with glass ionomer cement. The animals were sacrificed at 6 hours, 2, 4 or 7 days after treatment, biopsy samples were obtained, processed for immunostaining of COX-2 and viewed under light microscope. Data was statistically analyzed using Friedman and Kruskal-Wallis tests with significant level P<0.05. Results: Except in group I, COX-2 expression was decreased in all treatment group with the longer of observation time periods. In group II and group V, COX-2 expression was weaker than group III and IV at all periods time. However, there was no statistically significant differences of COX-2 expression among groups for each time period. Conclusion: The present study suggests that EEP and Ca(OH)2 were stronger than other materials test to suppressed COX-2 expression on inflamed rat dental pulp.

Keywords: Cyclooxygenase-2, Dental pulp, Inflammation, Propolis, Rat


Introduction

Propolis, or bee glue is a sticky, rubbery, brown, thermoplastic resin substance collected by bees from buds and exudates of the plants. Honey bees use propolis in their hives as a repairing crevice, and as a surface cover, hardener and preservative. Also, it is used as a repellent since it is applied inside the beehive that protect it from insects. There are a number of studies documenting the biological functions of propolis, its extracts and constituents. Several pharmacological activities have been described for propolis, including antibacterial, anti-inflammatory, accelerated wound healing and immunomodulatory. Among of its constituents, both flavonoid and non-flavonoid substances of propolis also showed anti-bacterial and anti-inflammatory activities. The chemical composition of propolis is very complex, depends on the collecting location, time and plant source. Commonly, the composition of propolis primarily consists of resinous (50%), wax (30%), essential and aromatics oils (10%), bee pollen (5%) and other substances (5%).

Recently, a new trend of using apitherapy has increased worldwide to treat many diseases. Using propolis in conservative dentistry and endodontic treatment to treat tooth and pulp diseases is a popular practice such as cariostatic agent in suppressing cariogenic bacteria, desensitizing agent to treat hypersensitivity dentin, intracanal irrigant, cavity disinfecting agent in atraumatic restorative treatment, medicament during root canal treatment, and also as direct pulp capping agent. Previous studies have demonstrated that propolis is toxic to dental pulp fibroblasts at 2 mg or above and not reduced the viability of dental pulp fibroblasts at 1 mg/mL.

Dental pulp is a loose connective tissue uniquely situated within the rigid encasement of mineralized dentin. However, dental pulp may become exposed due to caries, accidental mechanical during cavity preparation, tooth fracture or attrition and these can result an inflammation. Prostaglandins (PG) are important mediators of inflammation, the synthesis of which is initiated by release of arachidonic acid from cell membranes. Cyclooxygenases (COX) are enzymes responsible for conversion of arachidonic acids to PG. Two isoforms and one variant of COX (COX-1, COX-2, and COX-3, respectively) have been cloned and characterized. COX-1 is constitutively expressed in cells and is considered a housekeeping enzyme. The inflammatory and inducible effects of COX are mediated through COX-2, which is up-regulated in inflamed tissues.
and is at low or undetectable levels in healthy tissues. COX-2 has been recently shown to play an important role in the pathogenesis of dental pulpal inflammation. Trigona sp was one of honeybee species that we can found in South Sulawesi province, Indonesia. This honeybee species is stinging and can produce a lot of propolis. Recently, propolis been drawn much attention due to its anti-inflammatory properties. However, the potential anti-inflammatory effects of propolis on COX-2 in inflamed dental pulp tissue still remain to be elucidated. Therefore the aim of the present study was to analysis COX-2 expression on inflamed rat dental pulp after capped with propolis. The null hypothesis was that apply propolis on inflamed rat dental pulp could be decrease COX-2 expression.

Material and Methods

The experimental protocol was approved by the ethical committee of Faculty of Medicine, Hasanuddin University. The study was conducted at Bio-Pharmacal Laboratory, Faculty of Pharmacy and The Animal Research Development Center, Faculty of Veterinary Medicine, Gadjah Mada University, Yogyakarta and Department of Pathological Anatomy, Faculty of Medicine, Hasanuddin University, Makassar, Indonesia.

Two hundred and fifty grams raw propolis was obtained from Honey Bee Development Center, Hasanuddin University, Makassar which collected from honeycombs in Luwu regency, South Sulawesi Province, Indonesia. It was sliced and squashed with a mortal and pestle, and the extract was maseder for five days at temperature 45°C with continuous shaking with low rpm in a flask containing 1.25 L ethyl alcohol (ratio 1:5). Then filtered through Whatman paper No 1, and the extract was dried at temperature 60°C using a rotary evaporator to get Ethanolic Extract of Propolis (EEP). The residue was separated using toluene to get flavonoid and non-flavonoid fraction.

Eighty male Sprague Dawley rats (8-16 weeks old, weight 200-300 g) were used in this study. The animals were feed the standard food and water ad libitum throughout the study. The animals were feed the standard food and water ad libitum throughout the study. The room was maintain on a 12h light-dark schedule at a temperature of 26±2°C and a relative humidity of 60-70%. Rats were divided into five groups randomly, each consisting of 16 animals. Group I as a negative control was not conducted any treatment. Rats in group II, III, IV and V were anesthetized intramuscularly with ketamine (Ketalar®, Pfizer, Ireland) (65 mg kg⁻¹ body weight) and xylazine-HCl (Xyla®), Interchemie, Netherland) (7 mg kg⁻¹ body weight). Class I cavities (Black’s classification) were prepared on the occlusal surface of the permanent maxillary right first molar using a low-speed tapered round diamond bur (Intensiv®, Switzerland) (0.84 mm in diameter). The pulp was then exposed at the cavity floor using a dental explorer (Martin®, Germany) (0.35 mm in tip diameter) and allowed in the oral environment for 1 hour. After that, the exposed pulp directly capped with ethanolic extract propolis (EEP) (group II) (0.5 mg), flavonoid propolis (Group III) (0.5 mg), non-flavonoid propolis (group IV) (0.5 mg), or Ca(OH)₂ (Hydcal®, Technew, Rio de Janeiro, Brazil) as positive control (group V) (0.5 mg). Cavity was then air-dried and filled with permanent filling material (HS Posterior Extra®, GC, Tokyo, Japan).

In each group, 4 rats were sacrificed at 6 hours, 2 days, 4 days and 7 days respectively. The teeth and the surrounding bone were resected, fixed in Bouin’s fixative solution for 1 days, decalcified with acetic acid/formal saline for 7 days, embeded in parafin. Consecutive 6 μ thick sections were cut with bucco palatal direction using a rotary microtome, and collected on poly-L-lysine-coated slides. Endogenous peroxidase was blocked by incubation in 0.03% hydrogen peroxide in absolute methanol for 30 minutes at room temperature and washed with phosphate buffer saline. Immunohistochemistry was performed using a primary mouse monoclonal antibody against COX-2 (Neuromics, USA). After 30 minutes of incubation with block solution, sections were incubated with the primary antibodies at temperature 4°C overnight, followed by sequential 30 minutes incubations with biotinylated secondary antibody and streptavidin universal kit (ABC Elite Vectastain, Japan). The sections were counterstained with hematoxylin solution and mounted with glycerol-gelatin and viewed under a light microscopy (Leica®, Olympus BX41-U-CA, Tokyo, Japan).

The evaluation of COX-2 expression in dental pulp was carried out according to Faleiro-Rodrigues et al. based on two parameters: (1) The intensity of COX-2 expression (no detected=0; weak=1; moderate=2; strong=3) and (2) percentage of COX-2-positive cells (0%-0; 1%-10%=1; 11%-50%=2; 51%-100%=3). The sum of score from both parameter was then grading as following: Score 0=grade 0 (weak expression); Score 1-2=grade 1 (weak expression); Score 3-4=grade 2 (moderate expression); Score 5-6=grade 3 (strong expression). The scoring of COX-2 expression on rat dental pulp was blindly investigated by two separate pathologists. If there was any disagree-
**Table 1** The difference grade of COX-2 expression among time periods for each group

<table>
<thead>
<tr>
<th>Groups</th>
<th>Mean Rank</th>
<th>Friedman test</th>
<th>P</th>
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<tbody>
<tr>
<td></td>
<td>6 hours</td>
<td>2 days</td>
<td>4 days</td>
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<td>Without treatment</td>
<td>2.50</td>
<td>2.50</td>
<td>2.50</td>
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<tr>
<td>EEP</td>
<td>2.75</td>
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<td>2.63</td>
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<tr>
<td>Flavonoid</td>
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<td>2.85</td>
<td>2.80</td>
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<tr>
<td>Non-flavonoid</td>
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<td>Ca(OH)₂</td>
<td>2.75</td>
<td>2.75</td>
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Note: Significant difference at P<0.05

**Table 2** The difference grade of COX-2 expression among groups for each time period

<table>
<thead>
<tr>
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<th>Mean Rank</th>
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Note: Significant difference at P<0.05

ment between the evaluators, the sample under discussion was jointly analyzed until a consensus was reached. Statistical package for the social sciences (SPSS Inc. 1989-1999, Chicago, IL, USA) was used for statistical analysis. The Friedman test was used to analysis the significant difference of COX-2 expression between observation time periods for each group. Meanwhile, for analysis the significant difference of COX-2 expression between the groups for each observation time period were carried out by Kruskal-Wallis test. The significant statistical difference was accepted at P<0.05.

**Results**

Histogram of COX-2 in rats dental pulp tissue of all groups after 6 hours, 2 days, 4 days and 7 days of application can bee seen at figure 1. It showed that COX-2 expression was decreased in group II, III, IV and V with the longer observation time periods. Only weak expression of COX-2 showed in group I, II and V at all time periods. In contrast, in group III there was weak and moderate expression of COX-2 at all time periods, while in group IV showed strong expression of COX-2 at 6 hours, 2 days and 4 days. However, the results of Friedman and Kruskal-Wallis tests showed no significant differences (P>0.05) of COX-2 expression among 4 times periods of each group table 1 and among 5 groups of each time period table 2.

No necrotic pulp tissues were found in all animals of treatment groups throughout the study. The photomicrograph of COX-2 evaluation is presented in only by the section from all groups at 6 hours and 7 days figure 2.

**Discussion**

During tissue growth and inflammation, the level of PG is generated by COX-2. Inflammatory mediators such as interleukin-1 (IL-1), tumor necrosis factor-α (TNF-α), growth factor, lipopolysaccharide and tumor cells are stimulators of the COX-2 expression. Many pathological cytokines, such as IL-1 and TNF-α, stimulate COX-2 mRNA expression and PGE-2 production, which lead to tissue destruction. It has been reported that PGE-2 might be involved in pathogenesis of pulpal inflammation because implicated in many inflammation processes, such as vasodilation, increased vascular permeability, chemotaxis and pain. Therefore, it can be used as an indicator of the severity of pulpal inflammation. COX-2 participates in the pathogenesis of pulpal inflammation. Inflamed pulp, fibroblast and macrophages expressed COX-2, leading to PGE-2 production. Study by Holt, Hutchins and pileggi found the up-regulated of COX-2 on inflamed dental pulp and low or undetectable levels on healthy dental pulp. In the specimen of radicular cyst, COX-2 level was found also high.

Anti-inflammatory and immunomodulatory activities of propolis and its constituents have been study by a number of researchers. The result of present study demonstrated that the expression of COX-2 was decreased after EEP, flavonoid propolis, non-flavonoid propolis and Ca(OH)₂ application on inflamed dental pulp with the longer observation time periods. However, the COX-2 expression after EEP and Ca(OH)₂ application weaker than flavonoid propolis and non-flavonoid propolis application. Our previous study also found the suppression dental pulp inflammation by propolis.

Propolis was known to have anti-inflammatory, anti bacterial, anti oxidative, and immunomodulatory activity that permit inhibition of COX-2 expression, perhaps via suppression of cytokines production and macrophages-derived nitric oxide. Considering that propolis is a complex mixture, synergistic interactions between its compounds must also be considered as an important factor in its anti-inflammatory activity. The anti-inflammatory activity of propolis not only depends on the presences of flavonoids and caffeic acid...
and caffeic acid phenethyl ester (CAPE), but also by additive active compounds such as ferulic acid, (hydroxyl) cinnamic acid and diterpene derivates. Previously study by Rossi et al. found that EEP, flavonoid (galangin) and non-flavonoid (CAPE) propolis were effective to inhibit COX-2 activity in macrophages but not to other non-flavonoid propolis substances (caffeic, ferulic, cinnamic and chlorogenic acids).

Presently, Ca(OH)₂ was used in this study as material of control positive group because until now it was known as the most promising capping agent for direct pulp capping treatment to preserve tooth vitality in an exposed pulp cavity. The weak expression of COX-2 on inflamed rat dental pulp caused due to its alkanity. This alkanity actually has a beneficial effect on the injured dental pulp tissue, insofar as it causes mild irritation and stimulates the conjunctive tissue to defend and repair itself, initiating an mild inflammatory reaction to control and eliminate the irritating agent. Study by Nelson-Filho et al. found that Ca(OH)₂ induces a lesser degree of inflammatory infiltrate in the initial hours and inducing subsequent tissue repairing.

**Conclusion**

This present study showed that the application of all material test on inflamed rats dental pulp tissue decreased COX-2 expression along with the increase of the observation time period duration. However, EEP has better effect than flavonoid propolis, non-flavonoid propolis and Ca(OH)₂ to decreased COX-2 expression on inflamed rats dental pulp tissue. Therefore, the present results suggest that ethanolic extract of propolis (EEP) as well as Ca(OH)₂ suppressed COX-2 expression on inflamed rat dental pulp.

**Acknowledgment**

The authors wish to thank Prof. S Pramono, Ph.D (Faculty of Pharmacy, Gadjah Mada University, Yogyakarta, Indonesia) for his technical assistance in propolis preparation.

**Conflict of Interest**

The authors report no conflict of interest.

**References**


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