The correlation of consuming thymoquinone extract of nigella sativa to tooth socket re-epithelization on diabetic induced rats

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Abstract

Objective: The purpose of this research is to determine the effect of Tq to re-epithelization at the margin of tooth socket post extraction in diabetic rats.

Material and Methods: Used 27 rats were induced diabetic with streptozotocin (STZ) dose of 50 mg/kg body weight of rats which divided into 3 groups that has been stained with hematoxillin-eosin and cytokeratin 19. They are P1 treated with thymoquinone (Tq), P2 treated with metformin, and K without treatment. Each group consist of 9 rats were divided into 3 subgroups based on day of observation.

Results: showed there was an improvement of re epithelialization percentage in the Tq treatment group compared with P2 and K groups.

Conclusion: Thymoquinone extract may decrease blood glucose level and increasing re-epithelialization in margin of the tooth socket after extraction in diabetic rats.

Keywords: Tooth extraction, Diabetic, Re-epithelialization, Thymoquinone


Introduction

Diabetes Mellitus (DM) is a disorder metabolism that can be characterized by hiperglycemia.¹² Prevalence of DM in Indonesia increase every year, from 382 million in 2013 became 592 million in 2015 and the number of incident in woman is higher than the man.³

Hiperglycemia occur because there is a destruction in pancreatic cells or dysfunction of insulin or both of them.⁴ Hiperglycemia in the long term will lead glycation process to lipid and protein that have resulted AGEs increasing (Advanced Glycation End-products) and then AGEs products in the blood circulation interact with RAGEs (Receptor of Advanced Glycation End-products).⁵ Increasing number of AGEs and RAGEs interactions in circulation causes dysfunction of neutrophil, decreasing ability of transendehotelial migration, suppressing activity of phagocytic cells and decreasing the number of neutrophil infiltration. Those of them causes microvascular and macrovascular disorders.⁶-⁸

DM in long term make wound healing disorder. It should take attention for surgery treatment such as tooth extraction that can effect to prolong of healing time and make dry socket or alveolar osteitis.⁹,¹⁰

The preventing for that cases are minimalizing of post extraction complications through controlling of blood glucose concentration. Metformin is a synthesis drug that is often used for DM therapy but it has side effects like lactic acidosis until causing death.¹¹-¹³ WHO suggest to consume herbal medicine for DM therapy. Thymoquinone (Tq) extracts of black cumin have an anti DM that commonly used. Thymoquinone extracts considered have an effect to increase tooth socket re-epithelization process in diabetic induced rats.¹⁴,¹⁵

Based on literature above, Tq extract considers can decrease blood glucose level by repairing of β-cell pancreas, and protect AGEs formation in the beginning until at the end.¹² We also consider that Tq extract can decrease matrix metalloproteinase (MMPs) such as MMP-2 and MMP-9. The decrease of MMPs will increase re-epitelyzation process.¹³

We conducted an experimental study that aims to determine the effect of Tq extract from black cumin toward re-epithelization of marginal socket post extraction in diabetic induced rats. We hope this research can contribute to increase herb medicine field and development of using herb medicine in dentistry.
Material and Methods

The research was conducted after the approval of the ethics committee. The research used laboratory experimental with post control only group design which used total amount 27 rats who adapted along 1 week.

All samples had been fasted for 8 hours, before induced by streptozotocin (STZ) 50 mg/kg body weight (BW) in 0.1 M citrate acid buffer pH 4.5. Soon as possible after injected, all the samples have given glucose solution to protect from hypoglycaemic shock. After all samples injected by STZ, diabetic rat categorize if they have blood glucose level ≥ 250 mg/dL, then diabetic rat samples deviated into 3 groups (P1, P2 and K group), each groups contain 9 rats which for P1 group was treated with Tq extract 80 mg/kg BW that solved in olive oil, then for P2 group as positive control was treated with metformin 100mg/kg BW that solved with aquadest and K Group as negative control was treated with aquadest. P2 dan K group treated three times a day, while P1 group once time a day by intra gastric tube for seven days. Then under anesthetized by ketamin 0.15 ml/ 100 gr BW tooth extracted on left mandible first molar were done for all samples, using excavator and clamp. Tooth socket extended with diamond bur round end no. 1 using micromotor contra angle handpiece low speed with 5000 speed rpm for 2 second. All samples continuously treated with Tq, metformin and aquadest after extraction.

Three samples in each treated group decapitated at 3rd, 7th and 10th day after extraction. Left mandible was taken from insisive until posterior teeth, after that samples soaked in fixation solution used formaldehyde buffer 10% for 24 hour. Decalcification was done by formic acid 10% for 7th days. Tissue slide performed through Paraffin-embedded tissue block cutting using microtom 6 µm in thickness. Two slides made from each samples. The first slide stained by Hematoxyline Eosin (HE) and second slide stained by Immunohistochemistry (IHC) with Citokeratin-19 (CK-19).

All slides were observed by light microscope with 40x and 100x magnification. Measurement of re-epithelization length was done using CK-19 expression slide with the method as follow:*

\[
R = \frac{l_{EET}}{l_{Wound}}
\]

\[R : \text{Re-epithelization (\%)}
\]

\[l_{EET} : \text{Lenght of EET (Epidermal Epithelial Tongue)}
\]

\[l_{Wound} : \text{Lenght of wound gap}
\]

(Note: \(l_{EET}\) measured from epithelial tooth socket edges to the end of new epithelization. \(l_{wound}\) measured from epithelial tooth socket edges to mesio servical of second molars)

The Data will be ratio value and analyze using kolmogorov smirnov to know whereas the data is normal distribution and homogeneity determine using Levene test. If the data normal distribution and homogen, the test is performed by one way annova with \(p<0.05\). If the data is not normal distribution and not homogen, Kruskal-wallis test will used. Then all data continued to correlation regression test.

Results

Figure 1 shows Blood Glucose Level (BGL) in each group on 3rd, 7th and 10th day. The averages of BGL of Group P2 (as positive control) and group K (negative control) decreased from 3rd to 7th day, but they increased in 10th day (538 mg/dL and 575 mg/dL respectively). It was different with P1 group (Tq treatment) whereas BGL is little bit increased from 3rd to 7th day, but it was decrease clearly in 10th day (334.3 mg/dL). Nevertheless all samples are still hyperglycemic as long as investigation.

Based on histopathological appearance of each group with HE and CK-19 staining. re-epithelization on the tooth socket margin begin from mesial side grow and moved toward apical irregularly in a granulation tissue along mesial wall of tooth socket. Re-epithelization in 3rd day on K group was seen clearly than P1 and P2 figure 2.

Re-epithelization of K group became worse in 7th day and 10th day but it seen better progress in P1 group than K and P2 group figure 3 and figure 4.

The Measurement of re-epithelization length of tooth socket in each group showed any variations, but P1 group showed re-epithelization length is better than P2 and K groups. The re-epithelization in
P1 group was reached 74.65% in length while 38.56% and 25.93% in length on P2 and K groups on 10th day post extraction figure 5 but statistically not significant (p>0.05) between each group.

Based on linear correlation regression test between BGL and average percentage of re-epithelization showed weak correlation for P1 group with R square value was=0.277 (weak), meanwhile P2 and K groups were very weak (R=0.002 and 0.001).

Based on the results above, it was any different re-epithelization level among each group. In P1 group, the re-epithelization length showed a progress on 3rd day to 7th day and 7th day to 10th day. They are 56.61%, 53.05% and 74.65% respectively, also in P2 group, there was re-epithelization progress but it is until 7th day only (52.59%) then decreased on 10th day that have average 38.56%, meanwhile in K group showed re-epithelization length was good enough in 3rd day (48.30%), after that re-epithelization was decrease until 10th day which is the re-epithelization length only reached 25.93%.

The differences of re-epithelizations length was influenced by the differences of treatment. The increasing of the re-epithelization length in P1 group on 10th day was associated with decreasing of BGL after treated with Tq extract.

The given of Tq 80 mg/kg BW in rat could decrease BGL after 45 days consuming. While in P2 group and K group, re-epithelization length is not good enough perhaps BGL is still high above of normal level. The decrease of BGL will accelerate the process of re-epithelization on the margin of tooth socket post extraction.

**Discussion**

Consuming tymoquinone extract in P1 samples showed that have effect to decreases BGL figure 1. Decreasing of BGL made inflammatory cells migrate to tooth socket area easily to performed granulation tissue. Which it is needed to stimulate and accelerate socket re-epithelization. Decreasing BGL through any pathway; one of them is decreasing oxidative stress. A decrease of oxidative stress by Tq could protect pancreatic islet β-cells from damage and also can stimulate pancreatic β-cell regeneration therefore insulin can produce adequately. The other way decreases BGL by Tq is suppressing

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**Figure 2** Histopathological appearance on 3rd day post extraction (HE staining: A, C, E and CK-19 staining: B, D, F). Re-epithelization begin from mesial tooth socket toward apical in granulation tissue along mesial tooth socket (head arrow) (K group: A-B; P1 group: C-D; P2 group: E-F. RE is re-epithelization tongue; AB=alveolar bone; S is tooth socket; M2 is 2nd molar. Magnification 40x)

**Figure 3** Histopathological appearance on 7th day post extraction (HE staining: A, C, E and CK-19 staining: B, D, F). Elongation of Re-epithelization from mesial tooth socket toward apically (head arrow) seem clearly over alveolar bone in P1 group than K dan P2 group (K group: A-B; P1 group: C-D; P2 group: E-F. RE is re-epithelization tongue; AB=alveolar bone; S is tooth socket; M2 is 2nd molar. Magnification 40x)
of gluconeogenic enzyme which plays an important role in the process of gluconeogenesis such as phosphoenol pyruvatecarboxykinase (PEPCK), glucose 6 phosphate, fructose 1,6 bisphosphatase. This enzyme in diabetic patient had been increasing because of insulin dysfunction then lead hyperglycaemia.\textsuperscript{5,12}

The group that treated by metformin (P2), there was no visible a significant decreasing of BGL and tend fluctuate, even, increased on 10th day. That metformin with doses of 100 mg/kg BW for 3 times/day could decrease BGL in diabetic rats. The mechanism after consuming metformin will decrease mRNA that codes PEPCK enzyme, which has plays in initiate of gluconeogenesis by catalyzed decarboxylation and phosphorylation reaction to change oxaloacetate into Phosphoenolpyruvate.\textsuperscript{20}

In this research, there is no effect of consuming metformin (P2 group) maybe caused by cross reaction of metformin with food in rat stomach. Rat feeding in this research was ad libitum. if consuming metformin while the stomach was full it could be inhibit drug absorption to reach Concentration maximum (Cmax) in blood. The present of food in stomach will decrease absorption process of metformin then reduces 40% of Cmax in blood.\textsuperscript{21}

The correlation between BGL and re-epithelization in this animal study are not show linearity (the linearity week and very weak), it was appropriate with the theory that state before.\textsuperscript{9,10}

Theoretically, the elongation of re-epithelization in diabetic patient will increase if BGL decrease. The slowly of wound healing can be related to the high value of BGL that can trigger increasing the forming Reactive Oxygen Species (ROS) which will improve the signaling NADPH Oxidase. This circumstance causes the phosphorylation and degradation of inhibitor of NF-κβ that is Ikβ. Phosphorylation and degradation of Ikβ can causes release NF-kβ and than NF-kβ translocate into the nucleus cell, so that occurs an increasing of expression regulation of pro-inflammatory cytokines such as TNF-α and IL-1β. The present of TNF-α regulation causes decreasing regulation of TGF-β1 which is useful to increase the proliferation of keratinocytes cell (epithelial cells) on the injured area and at the same time decrease apoptosis cells of re-epithelization.\textsuperscript{22,23}

**Conclusion**

Based on the histopathological appearance and re-epithelization measurement, it seems that the healing tooth socket of P1 group is better than P2 and K groups. The healing of the socket in K group was seen irregular retapeg and it could not migrate perfectly to close the tooth socket. Maybe the re-epithelization obstacle in K group occurred
because of dry socket, so it did not have enough connective tissue and bones supporting inside of tooth socket.

In this research all tooth socket group still not closure completely, because the BGL still high above normal, but the group which treated with Tq extract (P1) tend have better tooth socket re-epithelization than P2 and K group. Further research is needed Tq research for longer periods until BGL normally and re-epithelization process is completely.

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Conflict of Interest

The authors report no conflict of interest in this research.

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