ABSTRACT

Purpose: To investigate the effect of Glycogen Synthase Kinase-3 (GSK3) antagonist on pulp tissue repair in rabbits. Materials and Methods: 24 adult healthy male rabbits were used in this study divided into three main groups: Group I (control): 2 rabbits with normal teeth without pulp exposure. Group II (TG): The exposed pulps of right posterior teeth of 22 rabbits were capped with KolSpon in association with 34 nM Tideglusib. Group III (MTA): The exposed pulps of left posterior teeth of 22 rabbits were capped with Mineral Trioxide Aggregate (MTA). The 2 rabbits with normal teeth and 6 rabbits with pulp capped posterior teeth were euthanized after one day. Their DNAs were extracted and tested for expression of Axin2 and β-actin by qPCR. 8 rabbits were euthanized after 4 weeks, and another 8 was euthanized after 6 weeks. The teeth were collected and prepared for immunohistochemical study using anti-β-catenin antibody. Results: The qPCR results showed the higher expression of Axin2 in the Tideglusib group than control group (without treatment) by about 5 folds and about 1.5 fold in the MTA group. The immunohistochemical results revealed the increase intensity in the Tideglusib group than MTA at four weeks with no significance difference (p>0.05). Whilst after 6 weeks the higher expression was in Tideglusib group with significance difference (p<0.05). Conclusion: The Tideglusib drug stimulates the expression of Axin2 transcription factor gene which is known to play a key role in regulation of odontoblast and osteoblast differentiation.

INTRODUCTION

The dentin-pulp complex is the inner part of the tooth underneath the surface enamel layer in the crown of the tooth and thin layer of cementum in its root. Dentin is the hard tissue portion of this complex and it is formed by odontoblasts [highly specialized mesenchymal
cells]. Odontoblasts are also responsible for the dentin repair along its life\(^{(1,2)}\).

When tooth subjected to infection [dental caries] or trauma, the pulp tissue can be exposed and become infected leading to the activation of the natural repair process. This process is depending on the severity of the injury. With milder injury, affected odontoblasts may survive and secrete a reactionary type of tertiary dentin to increase the barrier between the injury and the cells of the pulp. Sever injury can lead to local odontoblast death then the tertiary (reparative) dentin is formed by odontoblast-like cells which are differentiated from mesenchymal stem cells in the pulp tissue \(^{(3-5)}\).

Although large lesions, as deep caries cannot be compensated by natural repair process, current clinical tooth repair involving the usage of mineral aggregates which are used to restore the cavity in dentin that formed after removal of trauma or caries. This mineral aggregates remain in the prepared cavity and will not degrade, so normal mineral volume cannot be restored completely \(^{(6,7)}\).

The activation of Wnt/\(\beta\)-catenin signalling is the immediate response to the tissue damage and shows to be crucial for provoking the cellular-based repair in various tissues \(^{(8,9)}\). This Wnt pathway is considered to be very important in regulation of different cellular functions such as differentiation, proliferation, and survival. Moreover it plays crucial role in many physiological processes such as stem cell differentiation, wound healing, angiogenesis and embryonic development \(^{(9,10)}\).

The essential component of the Wnt/\(\beta\)-catenin signalling pathway is the \(\beta\)-catenin and considered to be the bottle-neck by which the main signals pass \(^{(11)}\). It regulates many of genes in different biological processes \(^{(12,13)}\). For instance, studies showed that the activation of \(\beta\)-catenin in embryogenesis resulting in ectopic dental tissue formation, while inactivation of \(\beta\)-catenin in developing tooth leading to arresting of its development at the bud stage \(^{(14,15)}\).

Other studies demonstrated that tooth repair and development have many similar molecular mechanisms \(^{(16-18)}\). Moreover, several studies showed that \(\beta\)-catenin can regulate adipocyte, chondrocyte and osteoblast differentiation of stem cells \(^{(19-21)}\).

The key cytoplasmic component of Wnt/\(\beta\)-catenin pathway is Glycogen synthase kinase 3 (GSK-3) enzyme that cause degradation of \(\beta\)-catenin and Axin2 by phosphorylation, this occurs in the absence of binding of Wnt ligand. On the other hand, the presence of this ligand will result in inhibition of GSK-3 activity leading to inhibition of \(\beta\)-catenin phosphorylation and attenuate its degradation. Unphosphorylated \(\beta\)-catenin then enter into the nucleus, where it interacts with the transcription factors [TCF/LEF1], this interaction leading to the activation the expression of WNT target gene includes Axin2 (Fig.1) \(^{(22-24)}\).

All the previous data confirmed that Axin2 expression after tooth damage is up-regulated by Wnt/\(\beta\)-catenin signalling \(^{(25)}\). For this reason, inhibition of Wnt signalling by agonists causing stimulation of tertiary dentin formation leading to restoration of the lost dentin following tooth preparation with new formed dentin.

Recent studies of the involvement of the GSK-3 in many diseases pathways leading to emerge it in the
pharmacological chemistry research field as a very interesting therapeutic target for the development of selective inhibitors as potential new drugs\cite{24,26-29}. So glycogen synthase kinases 3 (GSK3) inhibitors drug have been developed. It up-regulate Wnt activity in various experimental studies very efficiently. Two studies of Tideglusib (4-Benzyl-2-(naphthalen-1-yl)-[1,2,4] thiadiazolidine-3,5-dione), is in clinical trials phase for Alzheimer’s disease treatment\cite{30,31}.

Given all the previous data, the authors hypothesized that reparative dentin may formed by participation of β-catenin in odontoblastic differentiation. This study was directed to test the ability of the GSK3 inhibitors drug to activate Wnt signaling pathway and stimulate Axin2 gene expression which subsequently stimulates tertiary dentin following experimentally induced pulp exposure.

**MATERIAL AND METHODS**

Twenty four adult healthy male rabbits of age ranging from 3-4 months, mean weight (2.8 ± 0.4) kg were used in this study. The rabbits were housed in Faculty of Pharmacy, Al-Azhar University and maintained in a 12 hr. dark-light cycle, with humidity of 50 ±5% and at a temperature of 22±3°C and good ventilation. The rabbits were fed fresh hay, water, and fresh vegetables. The study was conducted according to the guidelines of the institution’s Animal Ethics Committee of Al-Azhar University.

**Steps of induced pulp exposure:**

The rabbits were anaesthetized by equal parts of katamin and xylazen subcutaneous injection\cite{29}. Pulp exposure was induced by a round carbide bur (FG ¼) attached to a high speed hand piece until the dentin was reached, then the pulp was penetrated using a 30G needle. The pulp exposure was performed on right and left mandibular first molars.

**Pulp capping procedures:**

After pulp exposure, the animals were divided randomly into two experimental groups (I & II) and control group:

**Experimental Groups: Consists of 22 rabbits with 44 teeth, where in each rabbit, the pulp exposures were performed on right and left mandibular first molars.**

- Group I: Pulp exposures were done on left mandibular first molars and capped with mineral trioxide aggregate (MTA) (n= 22 teeth).
- Group II: Pulp exposures were done on right mandibular first molars and capped with KolSpon in association with 34 nM Tideglusib dissolved and diluted in DMSO (dimethyl sulfoxide) (n= 22 teeth).

A layer of Glass Ionomer Cement was used as a seal over the capping material. Post-operatively, the rabbits were given analgesic injection.

**Control Group: Consists of 2 rabbits with normal teeth without pulp exposure.**

**Cytotoxicity test\cite{32}**

Rabbit’s pulp cells [cell Line] were obtained from American Type Culture Collection. It was supplied from cell culture department, VACsERA (Cairo, Egypt). These cells were cultured using Dulbecco’s Modified Eagle Medium [DMEM] (Invitrogen/Life Technologies) supplemented with 10% Fetal Bovine Serum (FBS) (Hyclone), 10 ug/ml of insulin (Sigma), and 1% penicillin-streptomycin.

A volume of 100µl complete growth medium was poured in a 96-well plate with cell density of 1.2 – 1.8 × 10,000 cells/well; then 100µl/well of the tested agent were added to the plate and incubated for 24 hours before the MTT assay. For determination of the cell metabolic activity, MTT [3-(4, 5 Di-methylthiazol-2-yl)-2, 5- diphenyltetrazolium bromide, Sigma] was added to the conditioned media and controls after 24 hrs. Then the resulted formazan product was dissolved in 200 µ l of dimethyl sulfoxide per well (DMSO, Sigma). A background absorbance of multiwell plates at 690nm was measured by a Spectrophotometer and subtract from the 450 nm measurement.
At the day of scarification, each animal was decapitated after being euthanized by a lethal dose of diethyl ether anaesthesia. Their right & left molar segments of the jaws were carefully dissected out. The lower first molars of animals from control and experimental groups that sacrificed after one day were extracted and kept in ice cold PBS then the DNA were extracted from its pulp to be used in the PCR. The specimens (from 4 &6 weeks animals) were then prepared for immunohistochemical examination.

Real-time Polymerase Chain Reaction (PCR)

PCR technique was used to quantify the expression of genes (Axin2) and (β-actin)\(^{(33)}\), as their expression was used to indicate the activation of Wnt pathway. mRNA isolation was carried out from the dental pulp that collected from rabbits without injury (control) and one day after injury using RNeasy extraction mini kit as recommended by the manufacturer instruction. Nanodrop was used to quantify the RNA and reverse transcribed into cDNA. Beta-actin was used as housekeeping gene (Forward-GGCTGTATTCCCCTCCATCG, Reverse- CACAGTTGGTAACAATGCTGT) and Axin2 was the read-out for Wnt pathway activity (Forward-TGACTCTCCTCCAGATCCCA, Reverse-TGCCCAACACTTAGGTGACA). The reactions were incubated in a real-time thermal detection system (ROTOR-GENE 6000 HRM Real Time Rotary Analyzer) at 95°C for 10 min for one cycle and then 95°C/(15 sec), 60°C/(60sec) for about 40 cycles. Measurements for each donor were conducted in triplicate.

Immunohistochemical study by using anti-β-catenin antibody to detect β-catenin in the pulp cells:

The specimens were routinely fixed, dehydrated, cleared and paraffin wax embedded. Paraffin tissue sections were deparaffinized and rehydrated then antigen retrieval was conducted. The sections then were incubated with blocking buffer for one hour, then incubated with the primary antibody, rabbit polyclonal anti-Beta-catenin Antibody were purchased from (Bethyl laboratories inc. Montgomery, tx,USA). Detection of bound antibodies was done by the EconoTek HRP Anti-Polyvalent (DAB) kit were purchased from (ScyTek Laboratories, Inc., Utah, USA) as recommended by the manufacturer instructions. The slides were stained with diaminobenzidine (DAB) chromogen to detect the reaction product (Brown colored of epitope)\(^{(34)}\).

Histomorphometric analysis:

The histomorphometric measurements were performed using “Optics Image Analyzer Computer System” the analytical evaluation was done through the use of image analysis software (Image J, 1.41a, NIH, USA). Colour thresholding was performed automatically to convert the positive staining into measurement units (pixels) produced by the image analyzer program. The quantitative measurement analysis of immunohistochemical stain depends on the gray value from dark to light stain reaction which is from 0 up to 10: (0-1—intense, 2-3—strong, 3-4—moderate, 4 up to 7—weak). The data collected from the image analyzer were statistically analyzed.

Statistical analysis:

Statistical analysis was done using the (SPSS 20; SPSS, Chicago, IL, USA). Paired t-tests and independent t-tests were performed to compare the grade intensity of B-Catenin at 4 weeks and 6 weeks mean scores between Control Group (MTA group) and Experimental Group (Tideglusib group). P-values ≤ 0.05 was considered significant.

RESULTS

A. MTT assay

MTT assay showed that the highest concentration of Tideglusib that was not cytotoxic to the pulp cells after 24 hours in culture was 34 uM (Table 1, Fig 2).
Table (1) *Results of cytotoxicity test of Tideglusib drug on dental pulp cells (DPS):*

<table>
<thead>
<tr>
<th>Serial</th>
<th>Sample code</th>
<th>Molecular weight</th>
<th>Cells</th>
<th>Results (IC50 Um)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>Tideglusib</td>
<td>334.39</td>
<td>DPC</td>
<td>34±1.26</td>
</tr>
</tbody>
</table>

![Graph](image)

**Figure (2) Linear graph of cytotoxicity assay results**

B. PCR results:

The qPCR results showed that the higher expression of Axin2 in the tooth treated with Tideglusib than control group (without treatment) by about 5 folds and about 1.5 fold in the tooth treated with MTA (Table 2, Fig 3).

C. Immunohistochemical results:

The immunohistochemical results of Tideglusib drug after 4 and 6 weeks showed expression of β-catenin in most odontoblasts as well as endothelial cell of blood vessels, where the 6 week group showed more intense reaction than that of the 4 weeks group. (Fig.4, 5). In MTA group, after 4 weeks the immunostaining showed diffuse moderate cytoplasmic reaction around the negative nuclei of odontoblasts (Fig.6). After 6 weeks, it was noted that there was a slight increase in β-catenin; moderate reaction was shown in odontoblasts with weak reaction in sub-odontoblast cells as well as in endothelial cells of blood capillaries (Fig.7).

![Graph](image)

**Table 2: Results of PCR**

<table>
<thead>
<tr>
<th>Sample data</th>
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</tr>
</thead>
<tbody>
<tr>
<td>Sample code</td>
<td>Axin 2 FLD</td>
</tr>
<tr>
<td>Serial number</td>
<td></td>
</tr>
<tr>
<td>1</td>
<td>Ref (MTA)</td>
</tr>
<tr>
<td>2</td>
<td>Test (Tideglusib drug)</td>
</tr>
<tr>
<td>3</td>
<td>Control (Rabbit tooth cells)</td>
</tr>
</tbody>
</table>

![Graph](image)

**Figure (3) Bar graph of expression of Axin2 gene in dental pulps of control, ref (MTA) and test (Tideglusib) groups.**

![Photomicrograph](image)

**Figure (4) Photomicrograph of Tideglusib drug group after 4 weeks showing granular positive immunoreaction for β-catenin in dental pulp cells and blood capillaries; D: Dentin; DP: Dental Pulp (DAB, Bar = 50 µm).**

![Photomicrograph](image)

**Figure (5) Photomicrograph of Tideglusib group after 6 weeks showing strong granular positive immunoreaction for β-catenin in odontoblasts, pulp cells, around blood capillaries and dentinal tubules (DAB, Bar = 50 µm).**
Figure (6) Photomicrograph of pulp tissue of MTA group after 4 weeks showing granular positive immunoreaction for β-catenin in odontoblasts, most pulp cells and endothelial cells (DAB, Bar = 50 µm).

Figure (7) Photomicrograph of MTA group after 6 weeks showing moderate diffuse positive reaction of β-catenin in odontoblastic cells and weak reaction in subodontoblastic cells. Note weak reaction in endothelial cells of few blood vessels (arrows) (DAB, Bar = 50 µm).

Statistical analysis:

The analysis of collected data showed a significant decrease of the positive staining of β-catenin between the Tideglusib group and MTA group. The highest mean value of β-catenin was recorded in Tideglusib group in both 4 and 6 weeks, with only significant difference between the groups in 6 weeks (Fig 8, 9).

DISCUSSION

The hallmark of healing process of dental pulp is the reparative dentin that formed after direct pulp capping with any dental material. The cellular-based repair stimulation in various tissues is the target of any invented drug and it’s proven that the Wnt/β-catenin signalling pathway is responded immediately to any tissue injury and it seems to be crucial for its repair (35-36). Glycogen synthase kinase 3 (GSK-3) is enzyme that plays a key role in the Wnt/β-catenin pathway signal transduction that in the Wnt ligand /receptor binding absence, phosphorylates Axin and β-catenin causing ubiquitination and degradation (37).
In this study, results clarified that the inhibition GSK3 by the specific inhibitors causes activation of the Wnt/β-catenin signalling axis leading to expression of definitive specific genes. Moreover the biological efficacy of Tideglusib drug as inhibitor of GSK-3 in pulp tissue was evaluated using a rabbit model of pulp exposure.

The animal model selected in the present study was rabbits because of the bigger size of their teeth than other rodents’ teeth, which make it more suitable for dental procedures (38). Moreover, many teeth can be used in one rabbit reducing the number of animals used for any experiment (39). As a positive control, MTA was chosen for this study since several studies revealed that MTA is better than conventional restorative materials in promoting the formation of reparative dentin (40, 41).

The current experiment, a period of 4 and 6 weeks was used to study the effect of capping materials on pulp tissues. We used the same time-interval used before to examine the effect of Tideglusib drug in other rodents (7).

The cytotoxic effects of the Tideglusib drug on rabbit’s pulp cells were investigated using the MTT assay. It is a reliable and sensitive colorimetric assay which measures cells viability (42). In this study, treatment of rabbit’s pulp cells with different doses of Tideglusib drug showed that the highest concentration of Tideglusib that was not cytotoxic to the dental pulp cells after 24 hr in culture was 34 uM. These results were found to be different with the results of other study (7), which reported that the safety dose is 50uM; we suggested that this difference could be due to the different animal model as he used rats in his study whereas we used rabbits.

In current study we used the immunohistochemistry technique to detect the β-catenin antibody in the pulp tissue as the immune staining is important to functional description and may reveal additional information, where conventional histology is insufficient (43).

The results of our study showed increase in the β-catenin expression rate after four weeks in both groups (TG & MTA) with insignificant difference (P-value equal 0.415) between them; this supposed to be due to the activated Wnt/β-Catenin pathway in both groups. The results are coinciding with other studies which pointed the activation of Wnt/β-Catenin signalling pathway after damage and during the healing process (22, 25, 44). Moreover, this finding was in according with another study, which found that odontoblast-like cells and dental pulp cells under the exposed area were β-catenin positive and during in vitro odontoblastic differentiation of dental pulp cells there was upregulation of β-catenin. These data suggested that β-catenin may participate in formation of reparative dentin and also might play an essential role in differentiation of odontoblasts (10, 45).

The immunohistochemical findings of the present study were consistent with another study which showed that the odontoblastic differentiation of stem cells derived from apical papilla may enhanced by activation of β-catenin (46). In addition previous experiments suggested that β-catenin may interact with many transcriptional co-factors forming DNA binding complex which regulates target genes transcription (47, 48). This supports the null hypothesis of that increasing the expression of Axin2 is a result of increasing the β-catenin in nucleus which is done by GSK-3 inhibitor enzyme.

Data recorded from two groups (TG & MTA) after 6 weeks revealed a decrease in the mean values of the MTA group with significant difference (P-value equal 0.000) when compared to TG group. The differences between the two groups might be displayed as a result of feedback inhibition of the Wnt/β-catenin in MTA group. This feedback inhibition was due to the overexpression of Axin2 whereas in the Tideglusib group, the drug irreversibly inhibits the GSK-3 enzyme so the cycle is not inhibited yet. This suggestion was in agreement with previous research, which found that Axin2 is a down regulator of β-catenin level by the
ability of the Axins to complex GSK3-β, β-catenin, and APC (49). Regarding the Tideglusib drug, another study showed that Tideglusib drug can irreversibly inhibit GSK-3 which explains the uncontrolled level of β-catenin (50).

In this study the polymerase chain reaction (PCR) technique was used to detect the expression of Axin2 gene in the pulp tissues of normal and treated teeth. This technique was used because it offered accurate and sensitive way to quantify the mRNA levels that is essential in biomedical researches (51).

The results of present study showed that higher expression of Axin2 in the tooth treated with Tideglusib than negative control group (without treatment) by about 5 folds and about 1.5 fold in the tooth treated with MTA (positive control group). These finding might be explained by the activation of the Wnt/β-Catenin signalling pathway by Tideglusib drug which caused entrancement of β-catenin to enter the nucleus and releasing this transcription genes (Axin2); this results was consistent with another previous study, which reported the high expression of Axin2 in relation with Tideglusib drug (7).

It is still unclear if Axin2 has a direct promotive role in mediating β-catenin in process of odontoblastic differentiation although this study found that β-catenin regulates Axin2 expression. Therefore, other studies should be carried out to eliminate the network of regulatory mechanism of β-catenin in odontoblastic differentiation of dental pulp cell during formation of reparative dentin.

CONCLUSIONS

From the previously mentioned results, the following could be concluded:

1. The Tideglusib drug causes activation of the Wnt/ B-Catenin signalling pathway and inhibits the degradation of β-catenin. B-catenin accumulates in the cytosol then translocates into the nucleus.

2. Small molecule GSK-3 agonists (Tideglusib drug) delivered via a biodegradable collagen sponge stimulates the expression of Axin2 transcription factor gene which is known to play a key role in regulation of odontoblast and osteoblast differentiation.

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