Research Article

p38 MAP Kinase-Mediated Odontogenic Differentiation of Dental Pulp Stem Cells

William Cheng1, Yessenia Valverde2, Nam-Seob Lee3, Hassan Marzban4, Seung H. Chung2* and Satish B. Alapati2*

1Department of Endodontics, University of Illinois at Chicago, Chicago, Illinois, USA
2Department of Oral Biology, University of Illinois at Chicago, Chicago, Illinois, USA
3Department of Anatomy, Konyung University College of Medicine, Daejeon, South Korea
4Department of Human Anatomy and Cell Science, University of Manitoba, Winnipeg, Canada

ARTICLE INFO

Article history:
Received 28 May, 2020
Accepted 10 June, 2020
Published 3 July, 2020

Keywords:
DPSC
p38 MAPK
hypoxia
differentiation
odontoblast

ABSTRACT

The multipotent nature of dental pulp stem cells (DPSCs) promises regenerative endodontic potentials. Alterations in microenvironment have been shown to control the differentiation phenotypes of DPSCs. Understanding the biological mechanisms and finding the optimal DPSC differentiation protocols are crucial for successful DPSC engineering strategies in pulp and dentin healing. The aim of this study is to identify the role of p38 mitogen-activated protein kinase (p38) under normal and oxygen-deprived conditions (2%) to reveal its effect on odontogenic DPSC differentiation. Human DPSCs were isolated from healthy molars and underwent odontogenic differentiation in regular and osteogenic media treated with SB203580, a p38 inhibitor, for 72 hours, and then swapped with osteogenic media for 21 days under hypoxic condition. Immunohistochemistry and PCR analysis for the various odontogenic differentiation genes and proteins were performed. Our PCR data demonstrate that p38 inhibition resulted in a significant upregulation in odontogenic gene expressions such as DMP-1, DSPP, RUNX, and OSX in normal conditions. Under hypoxia, this effect was reversed. These results were further supported by DSPP immunohistochemistry. The DSPP expression under hypoxia was significantly weaker compared to the control. Our results indicate that p38 represents a negative regulator of the odontogenic DPSC differentiation in normoxia. Under hypoxia, p38 exerts a positive function of DPSC differentiation. Taken together, we identified the p38 and oxygen level as crucial factors to control odontogenic DPSC differentiation providing their essential roles in designing for successful pulp-dentin complex engineering strategies.

© 2020 S.B. Alapati, S.H. Chung. Hosting by Science Repository. All rights reserved.

Introduction

In recent years in the regeneration field of dentistry, much attention has been given to stem cell biology as they demonstrate the remarkable translational capacity of regenerating dentin and damaged tooth [1, 2]. Dental pulp stem cells (DPSCs) can be acquired relatively with ease compared to other stem cell kinds such as embryonic stem cells (ESCs) and induced pluripotent stem cells. Several studies, including ours, have shown that human DPSCs can successfully differentiate into odontoblast-like cells under special culture conditions [3, 4]. These differentiated cells were able to produce dentin.

Changes in the cultural micro-environment have been shown to affect the phenotypes of stem cells. Extracellular environments, such as extracellular matrix, secreted factors and direct cell to cell contact, have both positive and negative roles in stem cell differentiation and cell fate determination [5, 6]. Oxygen is one of the vital factors and has a variety of functions on stem cell proliferation and differentiation [7-10]. The natural oxygen level in the body is remarkably low – approximately 3%
inside the tissue compared to routine 20% oxygen level in the lab tissue culture [11]. Thus, careful consideration of the micro-environmental oxygen level for stem cell engineering strategies needs to be addressed.

Mitogen-activated protein kinases (MAPks) are an evolutionarily conserved family of enzymes. They are serine/threonine kinases that connect cell-surface receptors to critical regulatory targets within the cell and are essential mediators of various physiological functions in normal as well as disease conditions [12]. p38 MAPks (p38) are essential mediators of various physiological functions in normal and disease conditions [12]. The p38 has been implicated in roles in tooth development and enamel production as well as in the periodontal tissue remodeling of orthodontic tooth movement [13, 14]. Furthermore, the p38 appears to be involved in dentin regulation. Dentin matrix proteins promoted bone marrow stromal cell differentiation through p38 signaling [15].

The role of p38 in the differentiation of DPSCs, especially under hypoxia, is unknown. Several studies have demonstrated that if the microenvironment is altered, stem cell fate will also be altered [16]. The present study focuses on the molecular signaling of stem cells to differentiate into the appropriate lineage. Our aim is to validate the role of p38-mediated odontogenic DPSC differentiation in the context of oxygen-deprived conditions. Our data will provide an important foundation for future DPSC engineering for dentin regeneration in the damaged tooth.

Methods

I Cell Cultures

DPSCs were collected from immature human healthy molars, and the isolated cells were cultured and expanded using the explant outgrown method [17]. Human third molars were extracted for orthodontics cases in the clinic (UIC protocol #20110129). We also used commercially available cells (Celprogen, #36086). The DPSCs have been evaluated in cultures with the STRO-1, a stem cell marker (IHC and FACS analysis). Our analysis confirms over 99% of cells (398/402, N=6) used for our study were DPSCs. The DPSCs were cultured using explant outgrowth method and will not be used beyond passage 5. DPSCs were cultured in regular media and osteogenic media and treated with p38 antagonist – SB203580 for 72 hours in regular media (regular growth media), and osteogenic media. DPSCs between 2nd and 4th passages were used throughout the study. All experimental protocols used for this study were in accordance with the guidelines according to the Institutional Animal Care and Use Policy and approved by the IRB Protocol Committee at the University of Illinois at Chicago.

II Quantitative PCR Analysis

Isolated DPSCs were cultured according to the differentiation protocol above in 6 wells plate with triplicates at 40,000 - 50,000 cells/well concentration for the treatment of the C5aR inhibitor W-54011. Total mRNA was extracted with 0.8 ml Trizol (Invitrogen, CA) and analysed using the Fisher Scientific NanoDrop 2000 device. The cDNA samples were analysed using the Applied Biosystems SYBR green reagent system according to the manufacture protocol. The list of primers and their sequences used are provided in (Table 1).

<table>
<thead>
<tr>
<th>Table 1: List of primers and their sequences used.</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>CDCC2 forward</strong></td>
</tr>
<tr>
<td><strong>CDCC2 Reverse</strong></td>
</tr>
<tr>
<td><strong>DMP1 forward</strong></td>
</tr>
<tr>
<td><strong>DMP1 reverse</strong></td>
</tr>
<tr>
<td><strong>P38 forward</strong></td>
</tr>
<tr>
<td><strong>P38 reverse</strong></td>
</tr>
<tr>
<td><strong>MAPK forward</strong></td>
</tr>
<tr>
<td><strong>MAPK reverse</strong></td>
</tr>
<tr>
<td><strong>BMP1 forward</strong></td>
</tr>
<tr>
<td><strong>BMP1 reverse</strong></td>
</tr>
<tr>
<td><strong>FGF forward</strong></td>
</tr>
<tr>
<td><strong>FGF reverse</strong></td>
</tr>
<tr>
<td><strong>DSPF forward</strong></td>
</tr>
<tr>
<td><strong>DSPF reverse</strong></td>
</tr>
<tr>
<td><strong>RUNX2 forward</strong></td>
</tr>
<tr>
<td><strong>RUNX2 reverse</strong></td>
</tr>
<tr>
<td><strong>OSX forward</strong></td>
</tr>
<tr>
<td><strong>OSX reverse</strong></td>
</tr>
<tr>
<td><strong>JNK forward</strong></td>
</tr>
<tr>
<td><strong>JNK reverse</strong></td>
</tr>
<tr>
<td><strong>GAPDH forward</strong></td>
</tr>
<tr>
<td><strong>GAPDH reverse</strong></td>
</tr>
</tbody>
</table>

III Alizarin Red Staining

Fourth passage DPSCs subject to their respective control or treatment were fixed with 10% paraformaldehyde for 15 minutes then washed three times with distilled water. 1 mL of Alizarin red dye was added for 20 minutes then washed four times with deionized water with gentle rocking to remove any additional Alizarin red. 1 mL of 10% acetic acid was added to prevent the cells from drying. The cells were then subject to visual inspection and imaging.

IV Immunohistochemistry

Differentiating and differentiated DPSCs were fixed and saturated. Then, cells were incubated for 4 hours with rabbit anti-p38 (1:250, Sigma) and mouse anti-DSPF (1:500, Sana Cruz) or their respective control isotypes. Finally, cells were treated for 40 minutes with a mix of Alexa Fluor-594 anti-mouse IgG, Alexa Fluor-488 anti-rabbit IgG (2 µg/mL) and/or DAPI (1 µg/mL). The coverslips were mounted, and images were taken using a Leica DMi6000 B microscope. Fluorescence staining was statistically determined by using the integrated density of each condition using ImageJ 1.49v software.

V Statistical Analysis

For quantification of histology and immunohistochemistry staining intensity, ImageJ was used. Fixed areas of 1mm x 1mm or 2mm x 2mm were selected to analyze the number of differentiated cells. The
statistical significance was demonstrated using the Student’s t-test (P < 0.05 was considered significant).

**Results**

I Odontogenic Lineage Cell Markers Such as DSPP, DMP-1 and RUNX2 are Expressed in the Differentiating and Differentiated Cells from DPSCs

To demonstrate specific in vitro role of p38 in DPSC odontogenic differentiation both in normal and oxygen-deprived conditions, isolated DPSCs were differentiated into odontoblast-like cells using p38 inhibitor SB203580 by our modified differentiation protocol. The human DPSCs were acquired from molars and cultured using regular growth and osteogenic medium for 24 days. A general p38 inhibitor SB203580 was treated from the differentiation day (D) 1 to D24 (for more details, see a timeline in Figure 1A). The homogeneous purity of the DPSC population was checked and validated in our previous reports [3, 4].

We next analysed the cell identity after full differentiation using several odontoblast lineage markers by real-time PCR and immunohistochemistry. The differentiated cells demonstrated odontoblast-like characteristics as they expressed odontogenic cell markers such as dentin matrix protein (DMP)-1, runt-related transcription factor 2 (RUNX2) and dentin sialophosphoprotein (DSPP) mRNA (Figure 2) and DSPP protein (Figure 1C). These are well-known odontoblast lineage-specific genes [18, 19]. Alizarin mineralization staining also confirmed a calcification (Figure 1B). Thus, our comprehensive odontogenic marker analysis demonstrated a successful odontogenic differentiation of DPSCs both in normal and oxygen-deprived conditions.

Figure 1: A schematic timeline representation of odontogenic DPSC differentiation and p38 is expressed in DPSC-derived differentiating cells. A) DPSCs were acquired from healthy molars using the explant outgrowth method. After reaching confluency, DPSCs were cultured and treated with regular and osteogenic media for 4 days. C5aR inhibitor SB203580 was added to the osteogenic media from the differentiation day 1 to day 24. The medium was replaced every 3 days. Alizarin red staining demonstrates a successful calcium mineralization, which is the odontoblast characteristics, both in the SB203580 treatment and control groups. Representative image of differentiating DPSCs at day (D) 4. B) Alizarin red staining shows a successful calcium compound detection both in the SB203580 treatment and control groups. C-E) Double immunofluorescence staining with anti-DSPP and p38 (green) antibodies demonstrated that the differentiating cells express both DSPP and p38 confirming both a successful odontogenic differentiation and p38 expression in the DPSCs-derived cells. Scale bar: 50 μm.

II Differentiating DPSCs express p38

We next identified the p38 expression in DPSCs-derived differentiating cells. Double fluorescence immunostaining with anti-DSPP and p38 antibodies demonstrated that the differentiating cells express both DSPP and p38 confirming both a successful odontogenic differentiation and p38 expression (Figures 1C-1E).

III p38 Inhibition Resulted in a Significant Upregulation in Odontogenic Differentiation Gene Expression

PCR gene expression analysis of p38, fibroblast growth factor (FGF), bone morphogenetic protein 1 (BMP1), DMP-1, RUNX2, Cell Division Cycle 42 (CDC42), c-Jun N-terminal kinases (JNK), ON in the SB203580 treatment group compared to control (as represented value ‘1’ of respective genes). The data demonstrate significant upregulation of odontoblast lineage genes such as DSPP, RUNX2 and DMP-1 in the W-54011 treatment group compared to control (Figure 2).
**p38 MAPK-Mediated Odontogenic DPSC Differentiation in Hypoxia**

**IV p38 Inhibition Under Hypoxia Resulted in a Significant Reduction in Odontogenic Differentiation Gene Expression**

Under hypoxia with the p38 inhibition, odontogenic lineage genes such as DMP-1 and DSPP expression is significantly decreased compared to control (without p38 inhibition) and normoxia groups (with p38 inhibition) (Figure 3).

**V DSPP Expression in the Differentiated Odontoblast-Like Cells Is Decreased with p38 Inhibition and This Effect Is Reversed in Hypoxia**

To further support gene expression data, we next performed the DSPP protein immunostaining with DPSCs-differentiated cells. In normal cells, without hypoxia induction and SB203580 treatment, the DDPP expression is evenly present in the cytoplasm of the differentiated cells (Figures 4A-4F). However, the DSPP expression is significantly upregulated in the SB203580 treatment group (Figures 4B & 4D), suggesting p38’s negative role in odontogenic DPSC differentiation. Consistent with the PCR data (Figure 3), this effect was completely reversed in the oxygen-deprived condition. DSPP expression is significantly downregulated in the differentiated cells under hypoxia (Figures 4C & 4G).

**VI Hypoxia Induction Has Little Effect on Odontoblast-Like Cell Proliferation**

To demonstrate whether the change of odontogenic gene expressions is due to the increased or decreased cell number during DPSC differentiation, we next examined the proliferation capacity of the differentiated cells. The number of DSPP-positive cells was counted and analysed in a fixed area of 1 X 1 mm² in the captured images. The number of the differentiated cells did not show a significant difference between the normoxia (1.23 ± 0.14, N=4) and hypoxia (1.04 ± 0.17, N=4, p<0.05) groups (Figure 4G). Our analysis demonstrates that p38 signaling and hypoxia induction have little effect on the odontogenic proliferation of DPSCs.
In this study, we first successfully differentiated our isolated DPSCs into odontoblast-like cells. Our results demonstrate that p38 constitutes a negative regulator of odontogenic DPSC differentiation as the p38 inhibition induces a significant upregulation of odontogenic lineage gene expression such as DMP1, RUNX2 and DSPP. The hypoxia induction showed opposite results. Our results confirm that the physical changes in microenvironment, such as hypoxia, are important factors in altering the differentiation properties of DPSCs.

Traditionally, immature necrotic teeth with apical periodontitis have been treated with apexification procedures using either long-term calcium hydroxide treatment or the MTA barrier method. A major disadvantage of apexification is that there is no potential for further development of the immature root end in terms of dentin thickness as well as root length. Additionally, long-term treatment with calcium hydroxide may predispose the already fragile immature root to fracture. In 2004, Banchs and Trope proposed a clinical protocol that is similar to the current AAE guidelines for revascularization based on observations of revascularization of reimplanted teeth, disinfection of the necrotic canal space, and induction of a blood clot into the canal [20]. Their study also found successful results. Today, regenerative endodontic therapy is recommended as an alternative treatment to the current AAE guidelines for revascularization based on maintaining undifferentiated stem cell status and stem cell differentiation/proliferation. The ESCs can maintain a longer period of undifferentiated status under low oxygen [9]. Human ESCs and MSC-derived cells showed more bone and cartilage differentiation in oxygen-

Figure 4: DSPP expression in the DPSC derived odontoblast-like cells in normoxia and hypoxia.

**A-F** Immunofluorescence double immunostaining with anti-DSPP (red) at 26 days of differentiation demonstrating the control (A, D), SB203580 treatment group in normoxia (B, E) and SB203580 treatment group under hypoxia (C, F). DMP-1 is expressed uniformly in the cytoplasm of the DPSCs-derived cells both in normal (A, D) and hypoxia group (B, E). p38 inhibition induced an upregulation of DSPP expression (B, E). However, the DSPP expression is significantly downregulated in the SB203580 treatment group in hypoxia (C, F).

**G** DSPP fluorescence intensity was quantified using image J software (Left) and the number of the cells between the normoxia and hypoxia were counted and calculated as the control value “1”. The SB203580 treatment group in hypoxia resulted in a downregulation of DSPP expression of the differentiated cells (blue: 1.56 ± 0.08, N=4 versus orange: 0.73 ± 0.06, N=5, *** p<0.005). The number of the cells did not show a meaningful difference between the normoxia (1.23 ± 0.14, N=4) and hypoxia (1.04 ± 0.17, N=4, p<0.05) groups.

**Discussion**

In this study, we first successfully differentiated our isolated DPSCs into odontoblast-like cells. Our results demonstrate that p38 constitutes a negative regulator of odontogenic DPSC differentiation as the p38 inhibition induces a significant upregulation of odontogenic lineage gene expression such as DMP1, RUNX2 and DSPP. The hypoxia induction showed opposite results. Our results confirm that the physical changes in microenvironment, such as hypoxia, are important factors in altering the differentiation properties of DPSCs.

The roles of p38 in several stem cell differentiation and proliferation have been studied. However, the results of p38 in osteogenic and chondrogenic differentiation from mesenchymal stem cells (MSCs) have been controversial. Both positive and negative roles of p38 in bone tissue differentiation have been reported [21-23]. It appears that the role of p38 in stem cell differentiation is complicated, and many factors, including the cell culture condition, timing and dose of inhibitor treatment, seem to play vital roles in MSCs differentiation.

The triad of tissue engineering includes stem cells, a physical scaffold and signaling molecules. But even with an appropriate source of stem cells, the differentiation lineage may not be appropriate due to an altered microenvironment. Once a canal becomes infected, the microenvironment is changed; and once the canal is disinfected with antimicrobials, the microenvironment is also changed. One example of this is that in the presence of LPS, stem cells from the apical papilla introduced into root canal segments were found to shift towards osteogenic phenotype as opposed to the odontogenic phenotype [24]. Many other authors have found that if the microenvironment is altered, stem cell fate will also be altered [5-11]. Thus, careful control of the microenvironment is vital to the success of regeneration strategies. The present study focuses on the molecular signaling of stem cells for their differentiation into the appropriate lineage.

There have been a number of studies revealing the oxygen’s role in maintaining undifferentiated stem cell status and stem cell differentiation/proliferation. The ESCs can maintain a longer period of undifferentiated status under low oxygen [9]. Human ESCs and MSC-derived cells showed more bone and cartilage differentiation in oxygen-
deprived conditions [7, 8]. There were also more bone cells and elevated markers of osteogenesis [7]. Human ESCs in low oxygen tension had increased ability to produce type I and II collagen and had different CD markers present on their cell surfaces [8]. Finally, hypoxia has been used to promote differentiation into various cell lines while also having an influence on the production of growth factors involved in angiogenesis [11].

Surprisingly, our study demonstrates that the hypoxia induction oppositely regulates differentiation gene expression compared to normal conditions. This observation is consistent with our pilot study that immune complement fragment C5a plays a positive role in odontogenic differentiation of DPSCs; however, this effect was reversed in oxygen-deprived condition. C5a receptor blocker induced a significant downregulation of odontogenic differentiation genes such as DMP1, ON, RUNX2, DSPP in normal condition, but under hypoxia, these gene expressions were significantly increased (Pasiewicz et al., unpublished observation). We do not understand the exact mechanism of this phenotype change. Further in detailed molecular pathway analysis is needed. Previous studies have demonstrated that hypoxia induction increases the proliferation of several types of stem cells [7-9, 25]. However, either p38 inhibition or hypoxia induction did not result in a meaningful change in DPSCs proliferation. It seems that several environmental factors, including the differentiation stage, stem cell kinds, degrees of hypoxia and culture conditions, are crucial factors in the differentiation phenotype determination.

In this study, we report that the p38 and oxygen levels are important regulators in the differentiation of DPSCs both in normal and hypoxia conditions. Controlling the microenvironment in stem cell differentiation is a vital strategy for successful dentin repair. Our data will provide an important basis for future DPSC-mediated pulp-dentin regeneration engineering.

Acknowledgement

This study was supported by the NIH/NIDCR grant (to S. Alapati grant no. DE019514).

Author Contributions

Cheng W, Valverde Y, Chung SH contributed to conception, design, data acquisition (immunohistochemistry), analysis, and interpretation, drafted and critically revised the manuscript. Lee NS, Marzbhan H contributed data acquisition and critically revised the manuscript. Alapati S provided DPSCs and human tooth sections for the study and also contributed to design, data analysis (DPSC differentiation), and interpretation the results. All authors reviewed and gave final approval.

Conflicts of Interest

None.

REFERENCES

1. Mabel M Cordeiro, Zhihong Dong, Tomaotsu Kaneko, Zhaocheng Zhang, Marta Miyazawa et al. (2008) Dental pulp tissue engineering with stem cells from exfoliated deciduous teeth. J Endod 34: 962-969. [Crossref]
15. Yan Yu, Lijuan Wang, Jinhua Yu, Gang Lei, Ming Yan et al. (2014) Dentin matrix proteins (DMPs) enhance differentiation of BMMSCs via ERK and P38 MAPK pathways. Cell Tissue Res 356: 171-182. [Crossref]


22. J Li, Z Zhao, J Liu, N Huang, D Long et al. (2010) MEK/ERK and p38 MAPK regulate chondrogenesis of rat bone marrow mesenchymal stem cells through delicate interaction with TGF-beta1/Smads pathway. *Cell Prolif* 43: 333-343. [Crossref]

