VPS4B mutation impairs the osteogenic differentiation of dental follicle cells derived from a patient with dentin dysplasia type I

Qiang Li1,2, Fangli Lu3, Tianxuan Chen4, Ke Zhang4, Yuping Lu1, Xiaocong Li5, Yingying Wang1, Ling Liu1, Qing Tian1, Fu Xiong6 and Dong Chen1

A splicing mutation in VPS4B can cause dentin dysplasia type I (DD-I), a hereditary autosomal-dominant disorder characterized by rootless teeth, the etiology of which is genetically heterogeneous. In our study, dental follicle cells (DFCs) were isolated and cultured from a patient with DD-I and compared with those from an age-matched, healthy control. In a previous study, this DD-I patient was confirmed to have a loss-of-function splicing mutation in VPS4B (IVS7 + 46C > G). The results from this study showed that the isolated DFCs were vimentin-positive and CK14-negative, indicating that the isolated cells were derived from the mesenchyme. DFCs harboring the VPS4B mutation had a significantly higher proliferation rate from day 3 to day 8 than control DFCs, indicating that VPS4B is involved in cell proliferation. The cells were then replenished with osteogenic medium to investigate how the VPS4B mutation affected osteogenic differentiation. Induction of osteogenesis, detected by alizarin red and alkaline phosphatase staining in vitro, was decreased in the DFCs from the DD-I patient compared to the control DFCs. Furthermore, we also found that the VPS4B mutation in the DD-I patient downregulated the expression of osteoblast-related genes, such as ALP, BSP, OCN, RUNX2, and their encoded proteins. These outcomes confirmed that the DD-I-associated VPS4B mutation could decrease the capacity of DFCs to differentiate during the mineralization process and may also impair physiological root formation and bone remodeling. This might provide valuable insights and implications for exploring the pathological mechanisms underlying DD-I root development.

International Journal of Oral Science (2020) 12:22; https://doi.org/10.1038/s41368-020-00088-z

INTRODUCTION

Dentin dysplasia type I (DD-I, OMIM: 125400) is a rare, autosomal-dominant, heritable, non-syndromic disorder characterized by obliterated pulp chambers, diminutive roots, and severe hypermobility of teeth.1-4 This is also often combined with frequent periapical radiolucencies in apparently sound teeth, both in the deciduous and permanent dentitions, with an estimated incidence of 1/100 000.5-7 DD-I increases the risk of early tooth loss and causes functional and esthetic disturbances. The initial information on DD-I mainly originated from reports of isolated cases owing to a low incidence. The genetic defects underlying DD-I were subsequently identified by genetic screening in affected families. To date, gene mutations in SMO2, VPS4B, and SSUH2 in three affected families from different countries have been identified, which strongly suggests that this disease is genetically heterogeneous.4,8-10 Despite major advancements in knowledge regarding molecular and cellular involvement in DD-I, the pathogenesis of this dysplasia remains undefined.

IVS7 + 46C > G, a splicing mutation that is genetically linked to DD-I in the extended Chinese family of this patient, was identified in the VPS4B gene.10 This gene is located on chromosome 18q21.33 and encodes a member of the AAA ATPase family.11 The VPS4B protein is an important component of the endosomal sorting complexes required for the transport (ESCRT) machinery12 and plays crucial roles in multiple cellular processes, including the formation of multivesicular bodies, virus budding,13 the abscission of cytokinesis,14 and degradation of various membrane receptors.15 However, the role of VPS4B in the development of other cell types, especially odontogenic cells, remains unclear. In our previous study, we demonstrated that the patient with affected teeth not only had dentin malformations but also had teardrop-shaped lacunae and a decreased organic content in their cementum.4,16 Additionally, it has been reported that the affected teeth are supported by insufficient alveolar bone, and the cementum is thin, sparse, or absent.17,18 These findings provide important evidence that VPS4B may cause imperfect cementogenesis and potentially affect the surrounding alveolar bone during mineralization development.

Interestingly, the dental follicle that originates from cranial neural crest cells, is a loose connective tissue that spherically surrounds the developing tooth germ in the early stages of development.19-22 This dental follicle is considered the top candidate for the origin of cementoblasts since it can create

---

1Department of Stomatology, the First Affiliated Hospital of Zhengzhou University, Zhengzhou, China; 2The Academy of Medical Sciences, Zhengzhou University, Zhengzhou, China; 3Department of Stomatology, Henan Provincial Hospital, Zhengzhou, China; 4Implant Dental Clinic, Zhengzhou, China; 5Department of Stomatology, Puyang People’s Hospital, Puyang, China and 6Department of Medical Genetics, School of Basic Medical Sciences, Southern Medical University, Guangzhou, China

Correspondence: Fu Xiong (xiongfu@smu.edu.cn) or Dong Chen (chendongfmmu@163.com)

Received: 15 February 2020 Revised: 2 June 2020 Accepted: 4 June 2020
Published online: 31 July 2020
cementum-like tissues without epithelial cells in vivo. Dental follicle cells (DFCs) reside in this ectomesenchymally derived, sac-like connective tissue. The normal differentiation of DFCs is essential for cementogenesis as well as surrounding alveolar bone development and formation. Many studies have reported that the differentiation of DFCs is always coordinated with root development. Moreover, DFCs are highly considered for the generation of biological tooth roots and for the regeneration of alveolar bone. It has been reported that rat DFCs form a tooth root when seeded on scaffolds of a treated dentin matrix and transplanted into an alveolar fossa microenvironment. Recent studies have focused on the characteristic and osteogenic differentiation of DFCs in vitro when induced with the appropriate growth factors. However, thus far, no data exist on the potential functional roles that DFCs may have during root development in DD-I. DFCs can be obtained from impacted third molars and have been shown to possess the capability of osteogenic differentiation in vitro when induced with the appropriate osteogenic medium. Furthermore, the VPS4B gene is one of the important contributors to root formation and is widely expressed in human tissues. Therefore, DFCs are a valuable tool to investigate osteogenic differentiation and to explore the mechanisms through which VPS4B affects the functions of these cells.

In our present study, we used DFCs as valuable tools to investigate differences in osteogenesis between a healthy individual and a DD-I patient, with the aim of determining the impact of a VPS4B mutation on the osteogenesis capacity of DFCs in DD-I, which has not been previously explored. These findings may contribute to the further understanding of the pathological mechanisms underlying DD-I root development.

RESULTS
Characterization and growth potential of DFCs in vitro
During clinical treatment, the DD-I patient with the VPS4B mutation had the impacted mandibular wisdom tooth removed. The third molar (at the root developing stage) was extracted and collected from both the DD-I patient and an age-matched healthy adult who underwent orthodontic treatment after informed consent was obtained (Fig. 1).

First, human dental follicles were harvested and subjected to enzymatic digestion to isolate DFCs. Only a small number of single dental follicle tissue cells attached to the plastic surface, and non-adherent cells were removed by changing the medium. We found that the adherent tissues were visible after being cultured for 24 h. Subsequently, a large number of primary passage cells around the tissue mass grew radially for nearly 5 days. At this time point, the cells presented a fusiform, spindle, or polygonal morphology but also had an excess of epithelial tissue (Fig. 2a–ii, v). From the 7th to 10th days, the cells converged and covered approximately 80% of the bottom of the culture flask. In addition, the cells from passages 3 and 4, which were isolated by plastic adherence, exhibited spindle or stellate shapes and were characterized by a typical fibroblast-like morphology (Fig. 2a–iii, vi).

Vimentin and cytokeratin 14 (CK14) immunofluorescence staining was first performed to verify the source of the cultured cells. The DFCs were vimentin-positive and CK14-negative, indicating that the cells were derived from the mesenchyme (Fig. 2b). The morphology of the cells was not obviously different; however, the subculture time differed between the two groups. In general, DFCs with the VPS4B mutation appeared after 3–4 days, while 5–6 days were required for the control DFCs, suggesting that the VPS4B mutant cells grew faster. The 3-(4,5-dimethylthiazol-2-yl)-2,5-diphenyltetrazolium bromide (MTT) assay showed that the VPS4B mutation significantly increased the proliferation rate of DFCs compared to that of the control cells from day 3 to day 8, indicating that VPS4B is involved in cell proliferation (Fig. 2c).

Osteogenic potential of DFCs in vitro
The course of osteoblast differentiation in DFCs was analyzed through the following experiments. The formation of mineralized nodule deposits over a period of 14, 21, and 28 days in culture was confirmed through alizarin red solution (ARS) staining. The intensity of the mineralization was enhanced with the increase in culture time. However, DFCs with the VPS4B mutation showed significantly smaller and fewer calcium nodules than the control DFCs at every time point (Fig. 3a, b). ALP staining results showed that the number of positive cells in both cell types increased with increasing mineralized induction time, while there was a lower number of positive cells in the mutated DFCs at each time point (Fig. 3c, d).

Changes in osteogenic genes and proteins
To determine whether the VPS4B mutation affected DFCs differentiation, we analyzed the changes in the levels of osteogenic-specific mRNA and protein markers in induced DFCs using real-time polymerase chain reaction (RT-PCR) and western blotting, respectively. We isolated total RNA to compare gene expression in DFCs after 0, 7, 14, and 21 days of mineralization induction in both groups (both at passage 3). We found notable differences in the gene expression of ALP, OCN, BSP, and RUNX2. The gene expression levels of ALP, OCN, BSP, and RUNX2 increased gradually with increasing osteogenic induction time in both groups. However, the expression of each gene in DFCs with the VPS4B mutation was significantly lower than that in control DFCs at each time point (Fig. 4a).

Western blotting showed that the osteogenic-specific proteins ALP, OCN, BSP, and RUNX2 were expressed from day 7, and the expression levels of these proteins increased gradually in the two groups. The protein expression in DD-I DFCs was significantly downregulated compared with that in the control DFCs at every time point (Fig. 4b, c).

DISCUSSION
DD-I is a rare, autosomal-dominant dental dysplasia disorder affecting both deciduous and permanent dentition, with an unclear etiology. It was first reported as “rootless teeth” in 1922 by Ballaschmiede, who found a family with six children manifesting with short, blunted roots and premature tooth loss. The estimated incidence of DD-I is 1/100,000, and thus,
knowledge of this disorder mainly originates from reports of isolated cases. Individuals with DD-I frequently present with tooth hypermobility combined with diminutive roots, aberrant dentin formation, obliterated pulp chambers, and numerous periodical radiolucencies in non-carious teeth. Therefore, the condition currently remains a clinical management dilemma and often leads to challenges in genetic counseling. Interestingly, various studies have reported the existence of stem cells and precursor cells in the dental follicles of humans, rats, and other species, which possess multi-lineage differentiation potential for cementoblasts, periodontal ligament cells, and osteoblasts. DFCs are known to have important functions during the formation of the cementum and root-surrounding alveolar bone. The osteogenic abilities of DFCs are crucial for this physiological process. Presently, most studies have focused on the characteristic and osteogenic differentiation of DFCs in different kinds of diseases, such as cleidocranial dysplasia, which is the failure of tooth eruption associated with parathyroid hormone-related peptide. However, no studies have compared DFCs between patients with DD-I and healthy individuals. Thus, to gain a better understanding of this disease, we explored the relationship between a VPS4B mutation and the osteogenesis mineralization abilities of DFCs. We designed this study to test the hypothesis that the VPS4B mutation affects osteogenic differentiation in DFCs, thereby disrupting the modulatory effects of DFCs on the differentiation of osteoblasts and consequently interfering with root and surrounding hard tissue formation and remodeling. In this study, we compared the proliferation and induced osteogenesis abilities of DFCs from a patient with DD-I with those from a healthy control.

Until now, few studies have compared cell proliferation between patients with DD-I and healthy individuals. One study performed proliferation assays using Cell Counting Kit-8, which indicated that human dental pulp stem cells transduced with an shRNA-Vps4b lentivirus proliferated slower than their counterparts transduced with a control lentivirus over 7 days. However, in our study, the DD-I DFCs growth rates were significantly increased compared to the control DFCs growth rates after 3 days of culture. The different proliferative capacity between hDPSCs and DFCs in DD-I may be related to the different sources of these cells. Further studies may be needed to better understand the mechanism underlying the effects of the VPS4B mutation on cell proliferation.

In this study, we examined the role of VPS4B in DFCs osteogenic differentiation. To this end, we performed ARS and ALP staining to monitor mineralization, and we examined the expression levels of the osteogenic differentiation markers ALP, BSP, OCN and RUNX2. ARS and ALP staining provided further evidence that VPS4B is involved in the osteogenic differentiation of DFCs and therefore may regulate root formation and bone remodeling during development. Consistent with the data from other studies, our results show that DFCs cultured in osteogenic induction medium increased the expression of these osteoblast-specific markers. In this study, we analyzed the mRNA expression of these four factors, and significant variations were found between DD-I DFCs and control DFCs. Western blotting analysis also
demonstrated the same trend in DFCs between the two subjects. According to the above results, we confirmed that the VPS4B mutation could increase the proliferation rate and significantly downregulate the expression of osteoblast-related genes and proteins in vitro, indicating a decreased osteogenic capacity in the patient with DD-I due to the VPS4B mutation.

In our study, the detailed molecular mechanisms underlying the functions of VPS4B during DFCs proliferation and osteogenic differentiation have yet to be fully elucidated. Notably, Wnt/β-catenin signaling is functionally significant for root odontogenesis and cementogenesis. One study showed that VPS4B acts as a regulator of hDPSC proliferation and differentiation via Wnt-β-catenin signaling. Moreover, it has been demonstrated that the overexpression or knockdown of VPS4B in human gingival fibroblasts markedly increased or decreased the expression of CHMP4B, Wnt5a, and β-catenin. The VPS4B-CHMP4B-Wnt5a-β-catenin regulatory pathway might regulate odontoblast differentiation and root formation. Thus, we speculated that VPS4B might regulate the osteogenic differentiation of DFCs through Wnt/β-catenin signaling, subsequently influencing root formation and bone remodeling. Future studies that test whether VPS4B affects other molecules or factors in the Wnt or other signaling pathways will undoubtedly improve our understanding of the molecular mechanisms underlying DD-I root development.

In addition, this study is a single-patient study with certain limitations, and the patient’s mutation of VPS4B may be different from variants of this gene in other patients with DD-I. Studies in more patients with DD-I and an individualized analysis of their characteristics could be performed to investigate this disorder further in the future. This would allow researchers to develop a more comprehensive understanding of this disorder with stronger evidence.

**MATERIAL AND METHODS**

**Cell isolation and culture**

This was an experimental study. The study protocol and patient consent were reviewed and approved by the Medical Ethics Committee of the First Affiliated Hospital of Zhengzhou University, China (2019-KY-104). The homolateral mandibular third molars, which had an immature root apex, were extracted and collected from both the DD-I patient and the age-matched healthy adult who underwent orthodontic treatment after informed consent was obtained. The attached dental follicles were gently separated from the tooth surface, and the dental papilla tissue was discarded. The surfaces of the tissues were thoroughly washed with phosphate-buffered saline (PBS; HyClone, China) and minced using a sterilized scalpel. The minced tissues were digested in a solution of 1 mg/ml collagenase type I (Bioleaf, Shanghai, China) for 1 h at 37 °C to obtain a cell suspension. The supernatant was discarded after the cell suspension was centrifuged at 12,000 r·min⁻¹ for 5 min. Thereafter, dental follicle explants were seeded into T-25 culture flasks, which contained 2 ml of an α-modification of Eagle’s medium (α-MEM; HyClone, China), 10% heat inactivated fetal bovine serum (FBS; Gibco, Grand Island, NY, USA), and 1% penicillin/streptomycin (HyClone, UT, USA); the cells were incubated at 37 °C and 5% CO₂ in a humidified atmosphere. After 24 h, the medium was replaced with fresh 10% α-MEM. After cells had adhered to the plastic dish surface, non-adherent cells were removed by changing the medium after overnight
incubation. Thereafter, the culture medium was changed once every 3 days, and cell morphology images of each group were captured via a phase-contrast inverted microscope (Sunny, Ningbo, China). At 80% confluence, cells were harvested with 0.25% trypsin and 0.02% EDTA (Dingguo Biotechnology, Beijing, China) and passaged in α-MEM containing 10% FBS for subsequent experiments. Cells from passages 2 to 5 were used for the experiments in this study.

For induction of in vitro osteogenic differentiation, cells that were digested with 0.25% trypsin were grown on 6-well plates at a density of 2 × 10⁴ cells per well in 10% α-MEM. At 80% confluence, cells were replenished with osteogenic medium consisting of α-MEM with 10% FBS, 1% penicillin/streptomycin, 10⁻⁶ mol·L⁻¹ dexamethasone, 50 μg·mL⁻¹ ascorbic acid, and 10 mmol·L⁻¹ β-glycerophosphate (Sigma-Aldrich, St. Louis, MO, USA). The cells were then cultured for a period of 28 days, and the medium was then changed every 3 days.

Identification of isolated cells
The purified DFCs were further identified using vimentin and CK14 immunofluorescence staining. A single-cell suspension was formed by digesting the cells with 0.25% trypsin and centrifuging them for 5 min at 12 000 r·min⁻¹. The cells were plated on coverslips at a density of 2 × 10⁴ cells per well in a 24-well plate. At 70%–80% confluency, cells were fixed with 4% paraformaldehyde (PFA; Dingguo Biotechnology) for 20 min. After being washed in PBS three times for 5 min, the cells were permeabilized with 0.25% Triton X-100 (Sigma-Aldrich) at 37 °C for 15 min. After being washed in PBS, cells were blocked with 1% normal goat serum (ZSG-BIO, Beijing, China) at 37 °C for 30 min and incubated overnight with rabbit anti-vimentin (Gibco) and rabbit anti-CK14 (Gibco) at 4 °C. After another PBS wash, the cells were incubated with goat anti-rabbit secondary antibodies (ZSG-BIO) at 37 °C for 1 h. Following another PBS wash, 4′,6-diamidino-2-phenylindole (DAPI; Invitrogen, NY, USA) was used to stain cell nuclei at 37 °C for 3 min; staining images were captured using a fluorescence microscope (Nikon, Tokyo, Japan).

Evaluation of cell proliferation
Growth profiles of cells cultured between days 1 to 8 from the two groups were measured by an MTT assay. Briefly, cells at passage 3 were seeded at a density of 1 000 cells per well in 96-well plates containing 10% α-MEM. At each time point, 20 μL of MTT solution (1 mg·mL⁻¹; Sigma-Aldrich) was added to each well; thereafter, all the plates were returned to standard tissue incubator conditions for an additional 4 h. After the removal of the MTT solution, 150 μL dimethyl sulfoxide (DMSO; Sigma-Aldrich) was added to each well and incubated while shaking in the dark for 15 min. The cell proliferation profiles were determined from the measurements obtained at a wavelength of 490 nm with a microplate reader (Shanghai Precision Instrument Co. Ltd., Shanghai, China). Experiments for all samples were repeated in triplicate.

Alizarin red staining
After 14, 21, and 28 days of induction, the presence of mineralized depositions was determined by staining with alizarin red. The cells were rinsed with PBS and fixed with 70% ice-cold ethanol. After being washed with distilled water, the cells were treated with ARS
Sodium dodecyl sulfate-polyacrylamide gel electrophoresis (SDS-PAGE) was used to separate cellular debris. The extracted proteins were separated by 10% polyacrylamide gel electrophoresis. The protein mixtures were centrifuged to remove fluoride (PVDF) membranes. The membranes were blocked for 2 h at room temperature using 5% non-fat milk (BD Biosciences, San Jose, CA). The membranes were incubated at 4 °C overnight with the following primary antibodies: anti-OCN (1:1 000; Boster Biological Technology, Wuhan, China), anti-BSP (1:1 000; Boster Biological Technology), anti-ALP (1:1000; Proteintech Group, Wuhan, China), anti-RUNX2 (1:1 000; Proteintech Group), and anti-GAPDH (1:1 000; Boster Biological Technology). Subsequently, the membranes were incubated with horseradish peroxidase (HRP)-conjugated secondary antibodies (ZSG-Bio) for 1 h at room temperature. The immunoreactive bands were visualized using an ECL chemiluminescence reagent (Solarbio). The relative intensities were analyzed and compared to their respective controls using ImageJ software (NIH, USA).

Competing interests: The authors declare no competing interests.

**Table 1. Sequences of the primers used for the real-time PCR**

| Primers                        | Abbreviation | Sequences (5’ to 3’) |
|-------------------------------|--------------|---------------------|
| Alkaline phosphatase          | ALP          | forward: GGACCATTCACGCTCTTACAC reverse: CTTTGTAGGCAAGGCCCCATTG |
| Osteocalcin                   | OCN          | forward: CCCAGGGCGCTACTCTATCATAA reverse: GTGCAAGCAAACCTGTCACAGTC |
| Bone sialoprotein             | BSP          | forward: CTTGCAAGGATTGATACAGGTTAG reverse: ACTGTCGCTGGTATGGCCTTG |
| Runt-related transcription factor 2 | RUNX2       | forward: GAGGCAACAAAGTTCTATCG reverse: GTGTCGTGGCCGATGATC |
| Glyceraldehyde 3-phosphate dehydrogenase | GAPDH       | forward: GCACCGTCAAGGCTGGAAC reverse: TGGTGAAGAGCCAGTGGA |

**Real-time PCR analysis**

Total RNA from cells was harvested after 0, 7, 14, and 21 days of osteogenesis induction using TRIzol Reagent (Invitrogen) and subjected to reverse transcription using the cDNA Reverse Transcription Kit (Takara, Dalian, China) in separate groups. RT-PCR was performed using SYBR Premix DimerEraser (Takara) on a LightCycler 480 (Roche, Indianapolis, IN, USA). The sequences of the related gene primers for RT-PCR are presented below (Table 1). The 2^ΔΔCt method was used to calculate the relative mRNA expression of each target gene with GAPDH as the reference. All samples were assayed in triplicate.

**ACKNOWLEDGEMENTS**

This work was supported by grants from the Science and Technology Research Program of Henan Province (22170124), the Medical Science and Technology Research Program of Henan Province (SBGJ2018038), the Science and Technology Program of Guangdong (2019A030317019), and the Medical Scientific Research Foundation of Guangdong Province (A2019232).

**AUTHOR CONTRIBUTIONS**

D.C. and F.X. conceived the idea of the study and the experimental design. Q.L. designed and validated the primer pairs used. Q.L., F.L., T.C., K.Z., Y.L., X.L., Y.W., and Q.T. conducted the experiments and analyzed the data. Q.L. and F.L. created the figures and tables. D.C., F.X., Q.L., F.L., and L.L. wrote the paper. All authors reviewed the paper.

**ADDITIONAL INFORMATION**

Western blotting was carried out as described previously. Total protein was extracted from the cultured cells using RIPA lysis buffer (Beyotime, Nanjing, China), which was supplemented with a protease inhibitor cocktail according to the manufacturer’s instructions. The protein mixtures were centrifuged to remove cellular debris. The extracted proteins were separated by 10% sodium dodecyl sulfate-polyacrylamide gel electrophoresis (SDS-PAGE) and electrotransferred to polyvinylidene fluoride (PVDF) membranes (Amersham, Little Chalfont, UK). The PVDF membranes were blocked for 2 h at room temperature using 5% non-fat milk (BD Biosciences, San Jose, CA). The membranes were incubated at 4 °C overnight with the following primary antibodies: anti-OCN (1:1 000; Boster Biological Technology, Wuhan, China), anti-BSP (1:1 000; Boster Biological Technology), anti-ALP (1:1000; Proteintech Group, Wuhan, China), anti-RUNX2 (1:1 000; Proteintech Group), and anti-GAPDH (1:1 000; Boster Biological Technology). Subsequently, the membranes were incubated with horseradish peroxidase (HRP)-conjugated secondary antibodies (ZSG-Bio) for 1 h at room temperature. The immunoreactive bands were visualized using an ECL chemiluminescence reagent (Solarbio). The relative intensities were analyzed and compared to their respective controls using ImageJ software (NIH, USA).

**Statistical analysis**

All results were obtained from triplicate samples. Data are presented as the means ± standard deviation and compared by Student’s t-test or one-way analysis of variance using the SPSS 21.0 software package (IBM Corporation, Armonk, NY, USA). P-values < 0.05 were considered to be statistically significant.

**CONCLUSION**

In summary, the present results confirmed that the DD-I-associated VPS4B mutation affected the osteogenic differentiation of DFCs and therefore may regulate root formation and bone remodeling during development. These findings provide valuable insights and implications for rootless teeth in patients with DD-I. Further studies are still needed to determine how VPS4B regulates the factors related to root formation and bone remodeling, as well as the intrinsic mechanism of these clinical manifestations in DD-I patients.
REFERENCES
1. Barron, M. J., McDonnell, S. T., Mackie, I. & Dixon, M. J. Hereditary dentine disorders: dentinogenesis imperfecta and dentine dysplasia. Orphanet J. Rare Dis. 3, 31 (2008).
2. Da Ros Goncalves, L. et al. Periodontal status of patients with dentin dysplasia type I: report of three cases within a family. J. Periodontol. 79, 1304–1311 (2008).
3. de La Dure-Molla, M., Philippe Fournier, B. & Berdal, A. Isolated dentinogenesis imperfecta and dentin dysplasia: revision of the classification. Eur. J. Hum. Genet. 23, 445–451 (2015).
4. Chen, D. et al. Dentin dysplasia type I-A dental disease with genetic heterogeneity. Oral. Dis. 25, 439–446 (2019).
5. Kalk, W. W., Batenburg, R. H. & Vissink, A. Dentin dysplasia type I: five cases within one family. Oral. Surg. Oral. Med. Oral. Pathol. Oral. Radiol. Endod. 86, 175–178 (1998).
6. Hart, P. S. & Hart, T. C. Disorders of human dentin. Cells Tissues Organs 186, 70–77 (2007).
7. Khandelwal, S., Gupta, D. & Likhyan, L. A case of dentin dysplasia with full mouth rehabilitation: a 3-year longitudinal study. Int. J. Clin. Pediatr. Dent. 7, 119–124 (2014).
8. Bloch-Zupan, A. et al. Homozygosity mapping and candidate prioritization identify mutations, missed by whole-exome sequencing, in SMOC2, causing major dental developmental defects. Am. J. Hum. Genet. 89, 773–781 (2011).
9. Xiong, F. et al. Mutation in SSUH2 causes autosomal-dominant dentin dysplasia type I. Hum. Mutat. 38, 95–104 (2017).
10. Yang, Q. et al. A splicing mutation in VPS4B causes dentin dysplasia I. J. Med. Genet. 53, 624–633 (2016).
11. Inoue, M. et al. Nucleotide-dependent conformational changes and assembly of the AAA ATPase SKD1/VPS4B. Traffic 9, 2180–2189 (2008).
12. Gan, X. & Gould, S. J. Identification of an inhibitory budding signal that blocks the release of HIV particles and exosome/microvesicle proteins. Mol. Biol. Cell 22, 817–830 (2011).
13. Watanabe, T. et al. Involvement of host cellular multivesicular body functions in hepatitis B virus budding. Proc. Natl Acad. Sci. USA 104, 10205–10210 (2007).
14. Morita, E. et al. Human ESCRT-III and VPS4 proteins are required for centrosome and spindle maintenance. Proc. Natl Acad. Sci. USA 107, 12889–12894 (2010).
15. Jiang, D. et al. High expression of vacuolar protein sorting 4B (VPS4B) is associated with accelerated cell proliferation and poor prognosis in human hepatocellular carcinoma. Pathol. Res. Pract. 211, 240–247 (2015).
16. Ye, X. et al. Dentin dysplasia type I: novel findings in deciduous and permanent teeth. BMC Oral. Health 15, 163 (2015).
17. Ozer, L., Karasu, H., Aras, K., Tokman, B. & Ersoy, E. Dentin dysplasia type I: report of a case and literature review. Eur. J. Hum. Genet. 16, 247–251 (2008).
18. Chen, J., Lan, Y., Baek, J. A., Gao, Y. & Jiang, R. Wnt/beta-catenin signaling plays an important role in odontoblastic differentiation of human dental pulp stem cells through the Wnt-beta-catenin signalling pathway. Artif. Cells Nanomed. Biotechnol. 47, 2575–2584 (2019).
19. Reizai-Rad, M. et al. Evaluation of bone regeneration potential of dentin follicle stem cells for treatment of craniofacial defects. Cytotherapy 17, 1572–1581 (2015).
20. Chen, J., Lan, Y., Baek, J. A., Gao, Y. & Jiang, R. Wnt/beta-catenin signaling plays an essential role in activation of odontogenic mesenchyme during early tooth development. Dev. Biol. 334, 174–185 (2009).
21. Yang, Q. et al. A splicing mutation in VPS4B causes dentin dysplasia I. J. Med. Genet. 53, 624–633 (2016).
22. Han, C. et al. Periapical follicle stem cell: a promising candidate for cementum/periodontal ligament regeneration and bio-root engineering. Stem Cells Dev. 19, 1405–1415 (2010).
23. Han, C. et al. Periapical follicle stem cell: a promising candidate for cementum/periodontal ligament regeneration and bio-root engineering. Stem Cells Dev. 19, 1405–1415 (2010).
24. Han, C. et al. Periapical follicle stem cell: a promising candidate for cementum/periodontal ligament regeneration and bio-root engineering. Stem Cells Dev. 19, 1405–1415 (2010).
25. Yoo, S., Pan, F., Prpic, V. & Wise, G. E. Differentiation of stem cells in the dental follicle. J. Dent. Res. 87, 767–771 (2008).
26. Sun, J. et al. IP3R2 regulates osteoclastic resorption in xenogeneic-treated dentin matrix-based scaffolds. Adv. Healthc. Mater. 6, https://doi.org/10.1002/adhm.201700127 (2017).