CIC-7 Deficiency Impairs Tooth Development and Eruption

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CLCN7 gene encodes the voltage gated chloride channel 7 (CIC-7) in humans. The mutations in CLCN7 have been associated with osteopetrosis in connection to the abnormal osteoclasts functions. Previously, we found that some osteopetrosis patients with CLCN7 mutations suffered from impacted teeth and root dysplasia. Here we set up two in vivo models under a normal or an osteoclast-poor environment to investigate how CIC-7 affects tooth development and tooth eruption. Firstly, chitosan-Clcn7-siRNA nanoparticles were injected around the first maxillary molar germ of newborn mice and caused the delay of tooth eruption and deformed tooth with root dysplasia. Secondly, E13.5 molar germs infected with Clcn7 shRNA lentivirus were transplanted under the kidney capsule and presented the abnormal changes in dentin structure, periodontal tissue and cementum. All these teeth changes have been reported in the patients with CLCN7 mutation. In vitro studies of ameloblasts, odontoblasts and dental follicle cells (DFCs) were conducted to explore the involved mechanism. We found that Clcn7 deficiency affect the differentiation of these cells, as well as the interaction between DFCs and osteoclasts through RANKL/OPG pathway. We conclude that CIC-7 may affect tooth development by directly targeting tooth cells, and regulate tooth eruption through DFC mediated osteoclast pathway.

CIC-7 is encoded by CLCN7 in humans and the mutations in the CLCN7 gene cause two different types of osteopetrosis: autosomal dominant osteopetrosis type II (OMIM166600) and autosomal recessive osteopetrosis type IV (OMIM611490)1. Previously we identified two Chinese osteopetrosis patients with CLCN7 gene mutations. Besides the identical bone phenotypes of osteopetrosis, the patients presented hypodontia, enamel dysplasia, malformed teeth, altered tooth eruption, and impacted teeth 2. The tooth germs were able to form but did not erupt in Clcn7−/− mice 3. A spontaneous autosomal recessive osteopetrosis mouse model with the mutant allele in Clcn7 locus displayed no tooth eruption or tooth root formation4. A recent study found no root was formed in the molars of 3-week old CIC-7 mutant mice5.

CIC-7 is expressed in the ruffled border of osteoclasts and dysfunctions of the CIC-7 affect osteoclast-mediated extracellular acidification, resulting in disturbed dissolution of bone inorganic matrix3,6. As we know, osteoclasts play an important role in tooth formation and eruption7,8. It is possible that the dysplastic tooth and the impact of tooth germ in osteopetrosis patients or in Clcn7−/− mice are caused by the dysfunctional osteoclasts.

CIC-7 has been also reported in other types of cells or tissues such as neurons9, gingival tissues10, bone tissues11,12, etc, suggesting that CIC-7 might have new functions in these cells or tissues. The locations of CIC-7 in ameloblast, odontoblast, dental papilla cells and dental follicle cells (DFCs)13,14 also raise one hypothesis, whether CIC-7 directly affect tooth development.

Here we set up two in vivo models with a normal and an osteoclast-poor environment to examine the effect of CIC-7 on the tooth development and eruption. In addition, we investigated the role of CIC-7 roles in the differentiation and biological functions of several tooth cells, such as ameloblasts, odontoblasts and DFCs.

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Results

Local Clcn7 silencing impairs tooth morphogenesis and tooth eruption. Comparing to the control (Fig. 1A,D,G,H), the enamel and dentin at day 7 post-injection became discontinuous and had uneven thickness in chitosan (CS)-Clcn7-siRNA group (Fig. 1B,C,E,F). The integrity and continuity of enamel was interrupted (E,F) and the disorganized odontoblasts were found (J, asterisk). In P17 CS-Cln7-siRNA group, the tooth was not fully erupted (N,T). The malformed tooth showed the abnormal cusps and short roots (N, black arrow; T) or disappeared root (N, asterisk). Abnormal and irregular fibrous tissues surrounded tooth instead of bone tissue (N, black triangle). The disorganized dentin and pulp cavity were observed (O,P, asterisk). The shape of the tooth crown was irregular and the root dentin was disrupted (S, black arrow) and uneven. The continuity and integrity of cervical loop and Hertwig’s epithelial root sheath was disappeared (S, black arrow). Figure (D,E) are partial enlargement of figure (A,B,G,I,K,N,Q,T), respectively. Scale bars = 200 μm. (U–W) Micro-CT scanning image of mouse head. U. Lingual view of the control side treated with CS-negative-siRNA nanoparticle. V Lingual view of the experimental side treated with CS-Clcn7-siRNA nanoparticle. W. Palatal view of the maxilla. The left first molar was impacted and lost its regular crown shape and cusps (white arrow). Cont = CS-negative-siRNA treated group; Clcn7 = CS-Clcn7-siRNA treated group. Scale bars = 2 mm.

CIC-7 deficiency disrupts morphogenesis of tooth in kidney capsule. After 3 days of lentivirus infection, obvious GFP fluorescence could be detected in the infected tooth germs, meanwhile, the Clicn7 mRNA level was decreased about 70% in the Clicn7-shRNA treated group, compared to the negative-shRNA group (P = 0.0001, data not shown). The teeth were well developed with clear profiles including normal cusps and roots in both blank control groups (Fig. 2A,D,G) and negative-shRNA group (Fig. 2B,E,H,K,M). The abnormal morphological alterations were found in Clicn7-shRNA group, such as short and/or crooked roots, abnormal cusps and apical foramen (Fig. 2C,I,N). Part of preodontin became thicker and the odontoblasts lost their regular column shape (Fig. 2F). The imposing characteristics of AZON staining in Clicn7-shRNA group were demonstrated as

Figure 1. Histological and micro-CT characteristics of tooth germs treated with CS-siRNA particles. (A–F,K–P) H&E staining; (G–J,Q–T) AZON staining. The molars were harvested and sectioned at day 7 (P7) (A–J) and day 17 (P17) (K–T) from mice treated by control CS-siRNA or CS-Clcn7-siRNA. The P7 molar treated with CS-Cln7-siRNA showed malformation of enamel and dentin (B,C,E,F). The integrity and continuity of enamel was interrupted (E,F) black arrow: enamel; black triangle: ameloblasts. (I,J). Dentin became disrupted and uneven (J, black arrow) and the boundary between dentin and predentin was indiscernible (F, asterisk), and the disorganized odontoblasts were found (J, asterisk). In P17 CS-Cln7-siRNA group, the tooth was not fully erupted (N,T). The malformed tooth showed the abnormal cusps and short roots (N, black arrow; T) or disappeared root (N, asterisk). Abnormal and irregular fibrous tissues surrounded tooth instead of bone tissue (N, black triangle). The disorganized dentin and pulp cavity were observed (O,P, asterisk). The shape of the tooth crown was irregular and the root dentin was disrupted (S, black arrow) and uneven. The continuity and integrity of cervical loop and Hertwig’s epithelial root sheath was disappeared (S, black arrow). Figure (D,E) are partial enlargement of figure (A,B,G,I,K,N,Q,T), respectively. Scale bars = 200 μm. (U–W) Micro-CT scanning image of mouse head. U. Lingual view of the control side treated with CS-negative-siRNA nanoparticle. V Lingual view of the experimental side treated with CS-Clcn7-siRNA nanoparticle. W. Palatal view of the maxilla. The left first molar was impacted and lost its regular crown shape and cusps (white arrow). Cont = CS-negative-siRNA treated group; Clcn7 = CS-Clcn7-siRNA treated group. Scale bars = 2 mm.
the disorderly arranged periodontal tissues. Large amount of fibroblasts replaced the well-organized periodontal ligament, cementum and alveolar bone (Fig. 2L) compared with the control groups (Fig. 2J,K).

Impact of ClC-7 on the expression of enamel and dentin matrix. The expression level of DSP was decreased in the group under the treatment of Clcn7-shRNA lentivirus infection (Fig. 3A~F). After lentivirus infection or siRNA transfection, the expression of Clcn7 in ameloblast LS8 and odontoblast-like MDPC-23 cells was downregulated. Subsequently the expression level of Enam was decreased in LS8 cell line (Fig. 3G), while Dspp was not obviously changed in MDPC-23 cells (Fig. 3H).

Impact of ClC-7 on the differentiation of DFCs. We confirmed the general mesenchymal characteristics of DFCs by positive immunocytochemical staining for anti-vimentin and negative staining for anti-pan cytokeratin (Supplementary Fig. 1).

The osteogenic induction with treated dentin matrix medium (TDMM) upregulated the mRNA expression of Alp, Bsp, Opn, Col1, and Tgfb1 (P < 0.01) with various ratios; while these induced upregulations of Alp, Bsp, Opn, and Tgfb1 were inhibited after Clcn7 gene level was lowered by Clcn7 shRNA lentivirus infection (Fig. 4A~C). However, the lowered Clcn7 level did not have any effect on calcified node formation with alizarin red staining (Supplementary Fig. 2A~H).

Effect of ClC-7 on osteoclast formation through DFCs. DFCs became round and lucent, and looked like multi-dendrites cells under the condition of DFCs/MNCs co-culture (Fig. 5A,B). The monocytes survived no more than three days in LS8 cells/MNCs co-culture system (Fig. 5C), but formed TRAP-positive multinucleated cells at day 7 post DFCs/MNCs co-culture, and lowered ClC-7 expression level resulted in less TRAP-positive multinucleated staining cells (Fig. 5D~F). These TRAP+ cells showed small and shallow pits on the dentin slices, and the numbers and area of resorption lacunae were decreased in ClC-7 deficiency group (Fig. 5G~H, Supplementary Fig. 3). qRT-PCR demonstrated the upregulation of Opg and downregulation of Rankl after Clcn7 gene was silenced in DFCs (Fig. 5I).

Discussion
Resorption of alveolar bones by osteoclasts is essential for tooth eruption. The osteopetrosis patients caused by osteoclasts dysfunction usually presented impacted and malformed teeth and the involved genes include RANKL, TCIRG1, etc. ClC-7 is highly expressed in the ruffled membrane of osteoclasts, and responsible for acidifying resorption lacuna. Osteoclasts isolated from ClC-7 deficient mice failed to resorb calcified bone, which might contribute to the tooth malformations and eruption abnormalities in osteopetrosis patients with CLCN7 mutation and Clcn7 gene knockout mice.
In the present study, we formulated siRNA with chitosan to protect siRNA from degradation, extended its in vivo effects and mimicked the gene knockout effect of Clcn7 around tooth germs. In the Clcn7-shRNA group, the expression of DSP was reduced compared to the other two groups. We chose the first upper molar of P1 mouse as the siRNA target site, because the cusps of P1 molar germ begin to form while the development of root has not started yet, which is a good time point to observe tooth eruption and root development. We found that CS-Clcn7 siRNA nanoparticles directly targeted on some tooth tissues or cells. The early responding tissues to Clcn7 siRNA were enamel and dentin, later in root and periodontal tissue. The local injection of CS-Clcn7 siRNA eventually resulted in impacted or delayed tooth eruption as we expected. We also tested the effect of local injection of chitosan formulated siRNA of Atp6v0a3, a gene involved in osteopetrosis, and found that...
Atp6v0a3 siRNA may also target tooth cells and cause tooth malformation and root dysplasia after 17 injection days (Supplementary Fig. 4). Thus the local injection of chitosan formulated siRNA of certain genes around tooth germ enables us to test a local gene silence or knock out effect during tooth development and this method should be widely used in the future.

Figure 4. CIC-7’s effect on the expression of different marker genes of DFCs. (A) Real time PCR results with Clcn7 or negative control shRNA lentivirus treatment. The RQ values in the Clcn7-shRNA group, which represents the mRNA level, of Clcn7, Bsp, Opn, Alp, Cap, Coll, Tgfb1 are 0.370 (0.161, P = 0.000), 0.416 (0.214, P = 0.042), 0.154 (0.031, P = 0.017), 0.733 (0.103, P = 0.046), 0.521 (0.126, P = 0.022), 1.313 (0.272, P = 0.351), 2.082 (0.760, P = 0.132), respectively, compared to the negative-shRNA group. B. Realtime PCR results of osteoblast-/cementoblast-related genes after 3 days’ treatment of TDMM. The RQ values of Bsp, Opn, Alp, Cap, Coll, Tgfb1 in the TDMM group, are 254.23 (69.484, P = 0.001), 94.074 (49.338, P = 0.033), 11.706 (2.898, P = 0.005), 0.978 (0.146, P = 0.819), 7.427 (2.672, P = 0.006), 6.855 (4.825, P = 0.007), respectively, compared to complete medium control group. (C) Realtime PCR results of osteoblast-/cementoblast-related genes after 3 days’ TDMM treatment as well as lentivirus treatment. The RQ values of Bsp, Opn, Alp, Cap, Coll, Tgfb1 in the Clcn7-shRNA group, are 0.615 (0.06, P = 0.008), 0.482 (0.017, P = 0.000), 0.688(0.042, P = 0.006), 0.855 (0.063, P = 0.058), 0.903 (0.031, P = 0.033), 0.536 (0.101, P = 0.015), respectively, compared to the negative-shRNA group. **P < 0.01, *P < 0.05. Data are presented as mean (SD, P value).

Atp6v0a3 siRNA may also target tooth cells and cause tooth malformation and root dysplasia after 17 injection days (Supplementary Fig. 4). Thus the local injection of chitosan formulated siRNA of certain genes around tooth germ enables us to test a local gene silence or knock out effect during tooth development and this method should be widely used in the future.
In order to identify the above tooth malformations caused by the effect of ClC-7 on osteoclasts or on tooth cells, the tooth germs infected with Clcn7 shRNA viruses were transplanted under kidney capsule which provides an osteoclasts-poor environment. The malformations in the transplanted tooth germs infected with Clcn7 shRNA viruses provided the direct evidence of ClC-7's contribution to the tooth development through tooth cells. Meanwhile the milder changes in the transplanted tooth germs infected with Clcn7 shRNA viruses comparing to that in situ CS-Clcn7 siRNA injection further confirmed that the contribution of ClC-7 deficient osteoclasts to the abnormal tooth phenotypes in situ.

ClC-7 deficiency caused the lower expressed DSP protein level in dentin in our study. Enamel and dentin changes had also been found in other osteopetrosis mouse models, such as Src(−/−) mice20. Comparing to the dentin, less effect was found on enamel of Src(−/−) mouse, which quite similar to our results. Thus ClC-7 may affect enamel and dentin development by regulating various key molecules of ameloblasts and odontoblasts.

We did not find the lowered Dspp mRNA level in MDPC-23 cell line, which might be because of the different responses of Dspp to lowered Clcn7 level in vitro, and the lower ΔCT value of Dspp in MDPC-23 cells.

A recent study in 3-week old ClC-7 mutant mice found that the molars did not form roots and the incisors were smaller than their age-matched controls. But the authors did not find obvious changes in the enamel of incisors and they regarded that ClC-7 deficiency might not significantly disrupt amelogenesis3. The different enamel's reaction to ClC-7 deficiency in the reference 3 and our data might be related with the position of investigated enamel. We observed enamel changes in molars, while Wen et al. studied the ultrastruture of enamel in incisors. The incisors and molars showed different root abnormalities in osteopetrosis patients with CLCN7 mutations2.

**Figure 5.** ClC-7's effect on the role of DFCs mediated osteoclastogenic differentiation. (A) Phase contrast images of DFCs and MNCs co-cultured for 7 days at low magnification (A) and high magnification (B), white triangles in A represented DFCs, white arrows in A represented MNCs gathering round the DFCs and white arrows in B represent cellular process of DFCs. (C) Phase contrast image of LS8 cells and MNCs after 3 days co-culture. (D) TRAP-positive cells on day 7 of DFCs/MNCs co-culture at low magnification. (E) Magnified region from D. Red arrows represent TRAP-positive multinucleated cells. (F) Comparison of the number of TRAP-positive multinucleated cell number per field vision. The Clcn7-shRNA group showed less TRAP-positive multinucleated staining cells than that of the negative-shRNA group and the control group (P = 0.001). (G,H) Dentine resorption area per lacunae (μm², n = 50). The dentin resorption lacunae formed by osteoclast-like cells in the Clcn7-shRNA group are reduced in amount and became smaller in area, when compared with the negative-shRNA group (Pnumber = 0.003, Parea = 0.000) and the control group (Pnumber = 0.011, Parea = 0.000). Bold horizontal bars indicate average values. The circle represents the outliers and the small asterisk represents the extreme values. **P < 0.01, *P < 0.05. Clcn7 = Clcn7-shRNA group; Neg = Negative-shRNA group; Cont = Control group. Scale bars (A–E) = 200μm. (I) Realtime PCR results of Opg and Rankl after knocking down Clcn7 in DFCs. The RQ value of Opg was upregulated to 3.789 (1.406, P = 0.000) and Rankl downregulated to 0.589 (0.185, P = 0.008), respectively. Data are presented as mean (SD, P value).
and Clcn7 gene knockout mice. On the other hand, the incisors were covered by enamel only in one side and the incisor growth is quite different from molar, especially during their root development. The deeper mechanism involved in the above different enamel changes under ClC-7 deficiency condition needs further exploring.

The deformed roots and abnormal periodontal tissues in ClC-7 deficient group switched our attentions to dental follicle cells. As the heterogeneous cell lineage, DFCs may differentiate into several types of cells, such as periodontal-type, cementoblastic-type and osteoblastic-type. The upregulations of osteoblast related genes such as Alp, Opn, and Bsp in DFCs with TDM induction confirmed the ability of DFCs to differentiate into osteoblasts; however, the above changes could be inhibited by Clcn7 gene deficiency. Thus ClC-7 might take part in the regulating of osteogenic differentiation of DFCs.

One interesting finding is that the mRNA level of Tgfb1 was downregulated in this osteogenic process after lowering the ClC-7 level. TGF-β1 has been reported to be involved in the regulation of tooth development. The release and activation of TGF-β1 stimulates the synthesis of various extracellular matrix proteins and inhibits their degradation which possibly results in tissue fibrosis. Our previous work found that the deficient CIC-5 and CIC-3 were related with the upregulation of TGF-β1. Here again we found TGF-β1 also being involved in the regulation of ClC-7 but with different mechanisms.

The interaction between DFCs and osteoclasts has also been studied previously. Early in eruption, the coronal part of dental follicle accumulates monocytes which may differentiate into osteoclasts. TRAP positive monocytes were present in the dental follicle prior to the onset of eruption and then declined in number during eruption. Here we found that DFCs/monocytes co-culture system could induce the monocytes to differentiate into osteoclasts. The co-cultures of human/rat DFCs and osteoclast precursors have been applied in contact or non-contact systems before. Due to the inhibition function of OPG secreted by DFC, both systems turned out a lower efficiency of TRAP+ cell formation and fewer and shallow resorption pits. Those TRAP+ cells in co-culture of DFC and monocytes only formed few and shallow pits in dentin slices, which might also be foreign body giant cells. Our contact co-culture of mouse DFCs and monocytes confirmed the above findings.

Meanwhile, we found that the lowered expression of ClC-7 in DFCs resulted in less TRAP positive multinucleated cells, less dentin resorption pits and smaller resorption area. Thus ClC-7 deficiency in DFCs might inhibit osteoclastogenesis.

RANK–RANKL–OPG signaling axis and downstream transcription factors play essential roles in the regulation of osteoclastogenesis. DFCs are assumed to secrete RANKL/M-CSF through a paracrine way, whilst OPG secreted by DFCs may inhibit osteoclastogenesis. Consistent with the previous data, we found the mRNA expression of Rankl and Opg in DFCs, and the lowered Clcn7 level downregulated Rankl and upregulated Opg, which cause the inhibition of osteoclastogenesis. We suggest that ClC-7 is involved in the process of osteoclastogenesis through DFCs mediated RANK–RANKL–OPG signaling pathway. Meanwhile, we admitted that the mechanisms by which osteoclast precursors interact with DFCs and differentiate into osteoclasts in vitro and in vivo are more complicated than we expected, and our findings indicated that multiple factors might take part in the interaction between DFCs and osteoclast precursors under ClC-7 deficiency conditions.

In conclusion, our experiments demonstrate that ClC-7 deficiency affects tooth development and eruption in several pathways. The dysfunction of ClC-7 may directly influence the formation and calcification of enamel, and impairs the differentiations of DFCs and DFCs mediated osteoclastogenesis through RANK–RANKL–OPG signaling pathway during tooth eruption. All the above changes contribute to the malformed teeth and altered tooth eruption.

Materials and Methods
Local delivery of CS-siRNA nanoparticles around tooth germ. siRNA duplex preparation. The small interfering RNA (siRNA) duplexes targeting the mouse Clcn7 gene with or without Cy5.5 fluorescence were designed and synthesized by GenePharma (Shanghai, China) according to GenBank™ (NM_011930.3). The sequence of Clcn7 siRNA is: sense, 5′-GAG GAG GAA AGA CGA UAC ATT -3′; antisense, 5′-UGA UUC GUC UUU CUC CCU CTT -3′. The sequence of Atp6v0a3 siRNA is: sense, 5′-GAA GGA CCC UUC UCU GAG A -3′; antisense, 5′-UCU CAG AGA AGG GUC CUU C -3′. The control siRNA (sense, 5′-UUC UCC GAA CGU GUC ACG UTT-3′; antisense, 5′-AGG UGA CAC GUU CGG AGA ATT-3′) was provided by GenePharma (Shanghai, China).

Preparation of CS-siRNA nanoparticle solution. Chitosan (CS) (100-300kDa, 93.37% deacetylation) (MP Biomedicals, US) was purified following the previously reported methods. The chitosan stock solution was prepared in acetate buffer (300 mM, pH 5.5) to a final concentration of 0.8 mg/mL, then 1 mL chitosan solution was mixed with 20 μl 100 μM siRNA solutions and stored at −20℃ till use.

Local delivery of CS-siRNA nanoparticles and in vivo fluorescence imaging. The newborn Balb/c mice (Experimental Animal Center of Fourth Military Medical University) were anesthetized through the inhalation of ether. The left and right maxillary first molar germ was chosen as the targeting sites of Clcn7 siRNA, Atp6v0a3 siRNA, negative siRNA, respectively. In the preliminary experiments, CS-Cy5.5-siRNA was used to test the suitable injection location and the dosages of injected nanoparticles. The distribution of fluorescence was observed by in vivo imaging system (IVIS Lumina II, USA). Then the non-fluorescence CS-siRNA nanoparticles solution were injected 1–2 mm below the eye toward the tooth germ direction using a microsyringe, and the depths of injection was 2.5 mm in newborn mice. For the long-term gene silencing experiments, two additional local CS-siRNA injections were added every two days till alveolar bones were too hard to perform injection. All procedures of animal work were approved by Fourth Military Medical University, and all experiments were performed in accordance with guidelines and regulations of Experimental Animal Center of Fourth Military Medical University.
Tooth germs kidney transplantation. pGCSIL-GFP lentiviral vector (GeneChem, China) was used to generate shRNA against Clcn7 gene with the same siRNA targeting sequences. The plasmids were co-transfected into HEK 293T cells with lentiviral packaging plasmids to generate Clcn7 shRNA lentivirus (Clcn7-shRNA group) or negative shRNA lentivirus (negative-shRNA group). The mandibular first molar germs dissected from E13.5 Balb/c mice were infected with Clcn7-shRNA lentivirus (5000MOI/tooth germ) with the medium containing Dulbecco Modified Eagle Medium (DMEM), 10% fetal bovine serum (FBS, Gibco, USA) and 5μg/ml polybrene (GeneChem, China). After 24 hours’ infection, the tooth germs were transferred to the transwell system, and the medium was switched to complete medium and cultured for about 18 hours. Then the tooth germs were transplanted into the renal capsule of adult male Balb/c mice. After 2 and 4 weeks of transplantation, micro CT scanning was performed to observe the tooth formation. The kidneys containing tooth germs were harvested for histological analysis and immunohistochemical staining after 4 weeks of transplantation.

Micro-CT scanning and analysis. Siemens Inveon system (Siemens, Germany) was used for the live animal scanning and three dimension images reconstruction. For CS-siRNA nanoparticles injection model, the heads of P17 mice were separated and scanned with the parameters as 80kV, 500 μA, 500 ms exposure time, and scan angle of 360°. For the tooth germ and kidney subcapsular transplantation, after 2 weeks and 4 weeks of the transplantation, the mice and the kidneys with tooth germs were scanned with the same parameters as above, respectively. The experiment group and the controls were subjected to the same radiation.

Histological analysis and immunocytochemical staining. All the tooth samples and the kidneys containing tooth germs were performed for hematoxylin & eosin staining and AZON staining, respectively. The sABC immunohistochemistry staining kit (Boster, Wuhan, China) was used for the immunohistochemical staining or immunocytochemical staining. Antibodies included mouse monoclonal IgG anti-DSP (kindly provided by Dr. Rong Zhang from Fourth Military Medical University), mouse monoclonal anti-vimentin antibody, mouse monoclonal anti-pan cytokeratin antibody (Abcam, UK).

Cell Culture. Cells and primary culture of mouse dental follicle cells (DFCs). Odontoblast MDPC-23 cell line (gift from Dr. C.T. Hanks and Dr. Jacques E. Nor, University of Michigan) and ameloblast LS8 cell line (kindly provided by Dr. Malcolm L. Snead in USC, Los Angeles, CA) were cultured in DMEM containing 10% FBS at 37 °C in a 5% CO2 humidified atmosphere. Using modified Wise's method31, the dental follicle tissues were dissected from the tooth germs of 3–5 postnatal days Balb/c mouse, cut into pieces and digested in a solution of 2 g/L type 1 collagenase for 1 hour at 37 °C. The cells were cultured in α-MEM containing 20% FBS, sodium pyruvate (0.1 g/l), streptomycin (100 μg/ml) and penicillin (100 U/ml). The 3rd passage cells were used for all the experiments.

Lentivirus infection of cells or siRNA transfection. Cells (LS8 cells, MDPC-23 cell, and DFCs) grew up to 30% to 40% confluence were transfected with Clcn7 siRNA or control siRNA using lipofectin 2000, or infected with Clcn7 (or control) shRNA lentivirus at an MOI of 100. The mRNA were harvested after 72 hours for qRT-PCR analysis.

Quantitative realtime polymerase chain reaction (qRT-PCR). Quantitative PCR was performed as previously described methods30. The specific primers for amplification of Gapdh, Clcn7, M-csf (macrophage colony stimulating factor), Trap (acid phosphatase 5, tartrate resistant), Ctsk (cathepsin K) and Rank (receptor activator of nuclear factor-κB), Rankl (receptor activator of nuclear factor-κB ligand), Opg (osteoprotegerin), Alp (alkaline phosphatase), Bsp (bone sialoprotein), Tgfβ1 (transforming growth factor-β1), Cap (cementum attachment protein), Ocn (osteopontin), and Col1 (collagen type I) are shown in Supplementary Table 1.

Osteogenic assay. Treated dentin matrix medium (TDMM) was prepared as the following methods: After being soaked in deionized water for 5 hours with an interval of 20 minutes ultrasonic treatment per hour, the TDMM were ground into powder with the treatment of liquid nitrogen, and sterilized with Co60 irradiation. TDMM was made with a ratio of 20 g dentin powder per 100 ml α-MEM containing 20% FBS, sodium pyruvate (0.1 g/l), streptomycin (100 μg/ml) and penicillin (100 U/ml). The 3rd passage cells were used for all the experiments.

Osteoclast function assay. Co-culture of DFCs and monocytes (MNCs). The monocytes were isolated using a Mouse Bone Marrow Monocyte Cell Isolation Kit (TBD science, China) from the femur and tibia of 6-week-old male Balb/c mice according to the manufacturer’s protocol. For the establishment of DFCs/MNCs co-culture system, cell density of DFCs and MNCs were adjusted to 105/ml and 106/ml, respectively. LS8 cells/MNCs co-culture, DFCs alone, MNCs alone were set as the controls.

TRAP staining and dentin resorption assay. After being co-cultured for 7 days, the DFCs/MNCs were stained for TRAP using Acid Phosphatase, Leucocyte (TRAP) Kit (Sigma, USA). The number of TRAP-positive multinuclear cells (>3 multinuclear) was counted and compared among Clcn7-shRNA group, negative-shRNA group and control group. After 14 days co-culture of DFCs/MNCs on dentin slices (4 mm × 4 mm × 200μm), the number of resorptive lacunae of each dentine slice was counted under the scanning electron microscopy (SEM). The areas of the lacunae were measured with NIH Image J software (NIH, USA) for at least 50 lacunae each group.

Statistical analysis. All the experiments were repeated at least three times and the data were presented as mean ± SD (standard deviation). Comparison between two different groups was performed using
Independent-Samples T Test. Comparison among three different groups were performed using a One-Way ANOVO, and multiple comparisons were performed using LSD test and SNK test when data meet the homogeneity of variance, otherwise Dunnett T3 test and Dunnett C test were performed. The difference was considered to be statistically significant when the P value was $<0.05$ or $0.01$.

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**Author Contributions**

X.D. conceived the idea; X.D., B.W. and H.H. conducted study; H.W., M.P., J.N., Y.Z., Y.Z., J.L. and Z.W. performed experiment; H.W., M.P., J.N., X.D. and S.G. discussed and wrote the manuscript; X.D., B.W., H.H., S.G. and R.Z. revised the manuscript. All authors contributed to the discussion and interpretation of the results as well as the preparation of the manuscript.

**Additional Information**

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