Smad1/5 signal transduction regulates the ameloblastic differentiation of induced pluripotent stem cells

Abstract: Induced pluripotent stem (iPS) cells could be induced into ameloblast-like cells by ameloblasts serum-free conditioned medium (ASF-CM), and bone morphogenetic proteins (BMPs) might be essential during the regulation of this process. The present study investigates the signal transduction that regulates the ameloblastic differentiation of iPS cells induced by ASF-CM. Mouse iPS cells were characterized and then cultured for 14 days in epithelial cell medium (control) or ASF-CM. Bone morphogenetic protein receptor II (BMPR-II) siRNA, inhibitor of Smad1/5 phosphorylation activated by activin receptor-like kinase (ALK) receptors, and inhibitors of mitogen-activated protein kinases (MAPKs) phosphorylation were used to treat the iPS cells in combination with ASF-CM. Real-time PCR, western blotting, and immunofluorescent staining were used to evaluate the expressions of ameloblast markers ameloblastin, enamelin, and cytokeratin-14. BMPR-II gene and protein levels increased markedly in ASF-CM-treated iPS cells compared with the controls, while the mRNA levels of Bmpr-Ia and Bmpr-Ib were similar between the ASF-CM and control groups. ASF-CM stimulation significantly increased the gene and protein expression of ameloblastin, enamelin, and cytokeratin-14. BMPR-II gene and protein levels increased markedly in ASF-CM-treated iPS cells compared with the controls, while the mRNA levels of Bmpr-Ia and Bmpr-Ib were similar between the ASF-CM and control groups. ASF-CM stimulation significantly increased the gene and protein expression of ameloblastin, enamelin, and cytokeratin-14, and phosphorylated SMAD1/5, p38 MAPK, and ERK1/2 MAPK compared with the controls. Knockdown of BMPR-II and inhibition of Smad1/5 phosphorylation both could significantly reverse the increased expression of ameloblastin, enamelin, and cytokeratin-14 induced by ASF-CM, while neither inhibition of p38 nor ERK1/2 phosphorylation had significant reversing effects. We conclude that smad1/5 signaling transduction, activated by ALK receptors, regulates the ameloblastic differentiation of iPS cells induced by ameloblast-conditioned medium.

Keywords: Induced Pluripotent Stem Cells; Bone Morphogenetic Protein Receptors, Type II.

Introduction

Stem-cell-based tissue-engineering approaches are being considered as a potential method to generate tooth tissues that closely match the physical and biomechanical properties of natural teeth. Many studies have reported that dental stem cells, such as dental pulp stem cells (DPSCs), periodontal ligament stem cells (PDLSCs), stem cells from exfoliated
deciduous teeth (SHEDs), or from the apical papilla (SCAPs), are multipotent and can form structures resembling tooth tissue. These dental stem cells have been applied successfully to generate dentin pulp and the periodontal ligament cement complexes. However, the shortage of a patient’s own dental stem cells limits the use of this approach.

Induced pluripotent stem (iPS) cells are considered as a new cell source for tooth regeneration, resulting from the rapid progress of iPS cells generation and the related technical platforms. Arakaki et al. developed a dental epithelial cell line from the cervical loop at the apical end of the lower incisors, and demonstrated that co-culturing with this epithelial cell line could induce iPS cells to adopt an ameloblast-like phenotype, such as displaying an epithelial cell-like morphology with positive expressions of the ameloblast markers ameloblastin, enamelin (ENAM), and cytokeratin-14 (CK14). However, Yoshida et al. demonstrated that mouse iPS cells could be induced into ameloblast-like cells more efficiently by culturing them with the conditioned medium of epithelial cell rests of Malassez (ERM) cells compared with co-culturing with ERM cells. Furthermore, Liu et al. showed recently that ameloblast serum-free conditioned medium (ASF-CM) could induce ameloblast-like cells from mouse iPS cells successfully, which was evident by the positive expression of ameloblast markers AMBN and CK14. They further found that culturing mouse iPS cells in ASF-CM supplemented with bone morphogenetic protein 4 (BMP4) promoted odontogenic differentiation. By contrast, culturing mouse iPS cells in ASF-CM supplemented with the BMP inhibitor noggin reversed the above effect. These results highlighted the promising future of tooth regeneration from iPS cells and indicated that growth factors from BMPs family could be essential during the regulation of this process. Therefore, the regulatory mechanism of the ameloblastic differentiation of iPS cells should be investigated further to promote the application of iPS in tooth regeneration.

BMPs, members of the transforming growth factor β (TGF-β) superfamily, are multi-functional cytokines that regulate the growth and differentiation of many cell types, and play a major role in early embryogenesis and in subsequent organogenesis. BMP signaling is mediated via the activation of combinations of type I and type II serine/threonine kinase receptors. These receptors’ signals have been reported to activate the canonical TGF-β signaling proteins termed SMAD proteins. However, recent studies have also confirmed that BMPs might also activate the mitogen-activated protein kinase (MAPK) pathways directly, including P38, ERK1/2, and JNK. In addition, previous studies have shown that BMPs are important regulators in dental morphogenesis, including the epithelial–mesenchymal interaction and epithelial bud development. These findings have generated much interest in the effects of BMPs on the differentiation of mouse iPS cells into the odontogenic lineage. However, the signal transduction pathways that are active during the ameloblastic differentiation of iPS cells remain unknown.

In the present study, we explored the mechanism of the ameloblastic differentiation of iPS cells induced by ameloblast-conditioned medium. A BMP receptor 2 (BMPR-2) short interfering RNA (siRNA) was used to investigate the involvement of BMPR-2 in this process. Furthermore, the involvement of SMAD1/5 and MAPKs signaling in the ameloblastic differentiation of iPS cells was also evaluated.

**Methodology**

**Culture and characterization of mouse iPS cells**

The mouse iPS cells (C5 cell line) were obtained from the South China Institute for Stem Cell Biology and Regenerative Medicine, Chinese Academy of Sciences. The iPS cells were cultured as previously described. Briefly, mouse embryo fibroblasts treated with mitomycin-C were used as the feeder cells for iPS cells, and Dulbecco’s modified Eagle’s medium (Gibco, Paisley, UK) supplemented with fetal bovine serum (Gibco), penicillin-streptomycin, β-mercaptoethanol (Sigma-Aldrich, St. Louis, USA), sodium pyruvate (Gibco), and leukemia inhibitory factor (Chemicon, Billerica, USA) was used as the culture medium. To
characterize the mouse iPS cells \((n = 6)\), the protein expression of the embryonic stem cell (ESC)-specific markers OCT4 and SSEA-4 and the differentiated ESC marker SSEA-1 were detected by immunofluorescent staining. All experimental procedures were performed according to institutional guidelines previously approved by the Ethics Committee of the First Affiliated Hospital of Xi’an Jiaotong University.

**Preparation of ameloblast serum-free conditioned medium (ASF-CM)**

ASF-CM was prepared as previously reported. Briefly, the mandibular incisors isolated from forty 7-day-old mice were used to isolate the ameloblasts (Supplementary Figure 1), and all mesenchymal cells were removed by repeated purification using trypsin (Gibco, Paisley, UK). Epithelial cell medium (Sciencell, Carlsbad, USA) was used to culture the purified cells, and was changed every 2 d. The supernatant was then collected, filtered, and stored at \(-80 \, ^\circ \text{C}\). In order to assure the reproducibility of the experiment, we have mixed all the supernatants together, and then used them as ASF-CM for the subsequent experiments. In this way, the active components in the conditioned media were the same among the experiments.

**RNA interference**

The mouse BMPR-II siRNA sequence was synthesized by Invitrogen (Invitrogen, Life Technologies, Grand Island, USA), and the siRNAs were transfected into cells as previously described. Briefly, the nucleotide target sequence of BMPR-II was GCA GTA CTA GTT CTA GCT TGC, and the sequence directed to the non-expressed mRNAs (ACG UGA CAC GUU CGG AGA ATT) was used as a control. \(1 \times 10^6\) iPS cells were seeded into each well of six-well plates. After 24 h of incubation, lipofectamine 2000 (Invitrogen, USA) was used to transfect the BMPR-II siRNAs or the controls into the iPS cells, following the manufacturer’s protocols \((n = 6)\). Twenty-four hours after transfection, the protein level of BMPR-II were determined by western blotting.

**In vitro differentiation of mouse iPS cells**

In the control group, the iPS cells were cultured in epithelial cell medium, which was used to create the ASF-CM. For the BMPR-II knockdown experiment, the iPS cells treated with BMPR-II siRNA or the control vector were cultured separately in ASF-CM or epithelial cell medium (Control) for 14 d \((n = 6)\). For the pathway inhibition experiment \((n = 6)\), the iPS cells were cultured in epithelial cell medium (Control), ASF-CM, or ASF-CM supplemented separately with 200 nM LDN-193189 (a selective SMAD1/5 inhibitor activated by activin receptor-like kinase (ALK) receptors; Selleck, Houston, USA), 10 \(\mu\)M SB203580 (a selective p38 inhibitor; Cell Signaling Technology, Danvers, USA), or 10 \(\mu\)M U0126 (a selective ERK1/2 inhibitor; Cell Signaling Technology). All the control and conditioned media were changed every 2 d during the 14-day experimental period.

**Immunofluorescent staining**

The iPS cells were fixed by 4% paraformaldehyde, and permeabilized by 0.1% Triton X-100 (Sigma-Aldrich, St. Louis, USA). To characterize the iPS cells, the specimens \((n = 6)\) were incubated overnight at 4°C with mouse anti-OCT4 (1:100), mouse anti-SSEA-4 (1:100) or mouse anti-SSEA-1 (1:100), and were then incubated with a Cy3-conjugated secondary antibody (1:100, Molecular Probes, Breda, Netherlands) for 1 h, and subsequently with 4’,6-diamidino-2-phenyl indole (DAPI) for 3 min at room temperature. To evaluate the ameloblastic differentiation of iPS cells \((n = 6)\), goat anti-ameloblastin (1:100), goat anti-enamelin (1:100), rabbit anti-CK14 (1:100) primary antibodies, and FITC-labeled secondary antibody (1:200; Chemicon, Billerica, USA) were used. All primary antibodies were purchased from the Santa Cruz Biotechnology (Dallas, USA). Samples \((n = 6)\) were examined under a confocal microscope (FV1000, Olympus, Japan). Five fields of view at 400× magnification were selected at random, and the percentage of the number of immune-positive cells to total cells in each field was counted and averaged for statistical analysis.

**Reverse transcription PCR (RT-PCR)**

Gene expressions of Ambn, ENAM, Ck14, dentine matrix protein-1 (Dmp-1), and dentine sialoprotein (Dsp) were detected as described previously. Briefly, total RNA from the iPS cells cultured under each
differentiation condition was extracted using Trizol (Thermo Fisher Scientific, Grand Island, USA). A 7500 real-time PCR machine (Thermo, USA) was used to analyze the expression of the target genes using glyceraldehyde 3-phosphate dehydrogenase (Gapdh) as the internal control. The results were calculated as the relative quantification compared to the control group. Primer sequences for the genes are shown in Table. Data were collected from six independent pooled samples (n = 6).

**Western blotting**

Total protein from the iPSCs (40 mg) was fractionated by sodium dodecyl sulfate-polyacrylamide gel electrophoresis (Bio-Rad, Martinez, USA) and transferred onto a nitrocellulose membrane. The membrane was blocked with 5% non-fat milk and incubated with primary antibodies against phospho-SMAD1/5 (1:800, 9516), glyceraldehyde-3-phosphate dehydrogenase (GAPDH, 1:1000, 2118), phospho-p38 (Thr180/Tyr182) (1:800, 451), total-p38 (1:1000, 9212), phospho-ERK1/2 (Thr202/Tyr204) (1:800, 4370), total-ERK1/2 (1:1000, 4695), phospho-JNK (1:800, 4688), and total-JNK (1:1000, 9252). All antibodies were obtained from Cell Signaling Technology (Danvers, USA). Signals were visualized by incubation with a horseradish peroxidase-conjugated secondary antibody (1:5000, Zhongshan Golden bridge Biotechnology, China) and enhanced chemiluminescence detection. The stained bands were scanned and quantified using a densitometer (Syngene Bioimaging System; Frederick, USA) and Scion Image software (Frederick). Protein expression levels were normalized against β-actin for each sample. Data were collected from three independent pooled samples (n = 3).

**Statistical analyses**

The measurement of cell number was performed blindly by two independent observers (ZG and QQ) using Photoshop CS 12.0 software (Adobe, USA) and there was a high level of agreement between the two observers (all r > 0.9), thus the mean of the two measurements was used for statistical analysis. All data were reported as mean ± standard deviation (SD) for each group. The Shapiro-Wilk test was used to assess the normality of the distribution of the data and Levene’s test was used to assess the homogeneity of variance, and the assumptions of parametric tests were fulfilled for all data. Therefore, the statistical significance among groups was evaluated by analysis of variance (ANOVA) with post hoc comparison between groups by Tukey’s test. P-values less than 0.05 were considered statistically significant in all cases.

**Results**

**Ameloblast-conditioned medium induced ameloblastic differentiation of iPSCs via BMPR-II**

The iPSC colonies were positive for embryonic stem cell (ESC)-specific markers OCT4 and SSEA-4, but were almost completely negative for differentiated ESC marker SSEA-1 (Figure 1A), indicating the ESC-like properties. To clarify the mechanism underlying the ameloblastic differentiation of iPSCs induced by ASF-CM, we cultured the iPSCs for 14 days in the control medium or ASF-CM, and the gene

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**Table.** Primers for quantitative real-time RT-PCR.

| Genes  | Sequence (Forward) | Sequence (Reverse) |
|--------|--------------------|--------------------|
| AMBN   | CCTGAGACAATGAGACAGTTGGG | CTTGGGTAAAAGCGGATGC |
| ENAM   | TCACTCCTCACCCCTCATCA  | CCACTTGGTTTCTCCTATTTCC |
| CK14   | CCTACTTCAAGACCATTGAGGAC | CATGCGCAAGGTCAACTCT |
| Dmp-1  | TCGGAGATGAGCCGATCAAAG | ATTGTTGTTGTAACATCA |
| Dsp    | CTCCGAGGCTTGAAGACATTGGA | GCTGACGTTCCGATGTGATAG |
| GAPDH  | CACATGGAGAGGCGGGGGGG | GACGGACACATGGGSGAG |
expression levels of different BMP receptors were analyzed. As shown in Figure 1B, Bmpr-II mRNA levels were markedly increased in ASF-CM-treated iPS cells compared with the controls (p = 0.013), while the mRNA levels of Bmpr-Ia and Bmpr-Ib were similar between the ASF-CM and control group (p > 0.05). A similar increase in BMPR-II protein levels was observed by western blotting analysis (p = 0.022, Figure 1C), indicating that ASF-CM could induce the expression of BMPR-II. To further confirm the involvement of BMPR-II in the ameloblastic differentiation of iPS cells, we pre-treated the iPS cells with a BMPR-II siRNA before culturing them in control medium or ASF-CM, and then detected the expression of ameloblast-specific genes (Ambn, ENAM and Ck14) and odontoblast-specific (Dmp-1 and Dsp) genes. The transfection efficiency of BMPR-II siRNA is about 82%, and the protein level of BMPR-II was reduced dramatically by the BMPR-II siRNA (p < 0.05, Figure 1C). The PCR results showed that ASF-CM stimulation significantly increased the gene expression of Ambn, ENAM, and Ck14 compared with that of the control, while BMPR-II siRNA obviously attenuated the above ASF-CM-induced effects (all p < 0.05, Figure 2). However, the gene expression of Dmp-1 and Dsp was similar among the control, ASF-CM,
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Figure 2. RT-PCR analysis of the expression of ameloblast-specific genes (Ambn, ENAM, and Ck14) and odontoblast-specific genes (Dmp-1 and Dsp) in iPS cells induced by the control medium, ASF-CM, or ASF-CM+BMPR-II siRNA.

A

B

*p < 0.05, vs. control group. **p < 0.01, vs. control group. ***p < 0.01, vs. ASF-CM group.

Figure 3. Immunofluorescent staining and quantification of the expression of AMBN, ENAM, and CK14 in iPS cells induced by the control medium, ASF-CM, or ASF-CM+BMPR-II siRNA. Scale bar represents 50 μm.

*p < 0.01, vs. control group. **p < 0.01, vs. ASF-CM group.
and ASF-CM+BMPR-II siRNA groups (p > 0.05, Figure 2). Immunofluorescent staining showed that the iPS cells cultured in control medium were almost completely negative for AMBN, ENAM, and CK14 expression, while those cultured in ASF-CM showed strong cytoplasmic staining for AMBN, ENAM, and CK14 (Figure 3). Interestingly, the ASF-CM induced expression of AMBN, ENAM, and CK14 was also attenuated by the BMPR-II siRNA (all p < 0.05, Figure 3). However, the expressions of DMP-1 and DSP were not detected in the above three groups (data not shown). The quantitative data showed that ASF-CM stimulation increased the protein levels of AMBN, ENAM, and CK14 significantly compared with those in the control, while the BMPR-II siRNA attenuated the above ASF-CM-induced effects significantly (all p < 0.05, Figure 3).

**Ameloblast-conditioned medium activated SMAD1/5, P38, and ERK1/2 signal transduction during ameloblastic differentiation of iPS cells**

To investigate the molecular mechanisms during ameloblastic differentiation of iPS cells via BMPR-II activation, the phosphorylation of SMAD1/5 and MAPKs pathways were detected. The result showed that ASF-CM stimulation increased the phosphorylation of SMAD1/5, P38, and ERK1/2 significantly compared with the control (all p < 0.05, Figure 4); however, the phosphorylation of p-JNK in the ASF-CM stimulation group and the control group were similar (all p > 0.05; Figure 4).

**SMAD1/5 phosphorylation, but not P38 or ERK1/2 phosphorylation, is involved in the ameloblastic differentiation of iPS cells induced by the ameloblast-conditioned medium**

To investigate whether the SMAD1/5 and/or the P38 or ERK1/2 pathways are involved in the ameloblastic differentiation of iPS induced by the ameloblast-conditioned medium, inhibitors of SMAD1/5, P38 or ERK1/2 phosphorylation were used to stimulate iPS cells in combination with ASF-CM treatment. The results showed that SMAD1/5 inhibitor LDN-193189 decreased substantially the gene and protein expression of AMBN, ENAM, and CK14 induced by ASF-CM (all

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**Figure 4.** Western blotting analysis and quantification of the phosphorylation level of SMAD1/5 and P38, ERK1/2, and JNK MAPKs pathways in iPS cells induced by control medium or ASF-CM. **p < 0.01, vs. control group.**
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p < 0.05, Figures 5 and 6), while neither the P38 nor ERK1/2 inhibitors had any blocking effect on the ASF-CM-induced increase in the expressions of AMBN, ENAM and CK14 at the gene or protein level (all p > 0.05, Figure 5 and 6).

Discussion

Recent study showed that BMP4 is an essential growth factor for the effective induction of ameloblastic differentiation of iPS cells. In the present study, we showed, for the first time, that inhibition of BMPR-II, one of the major receptors of BMP4, could block the ameloblastic differentiation of iPS cells induced by ameloblast-conditioned medium, which was characterized by decreased gene and protein expression of ameloblast markers AMBN, ENAM, and CK14. Furthermore, although the ameloblast-conditioned medium could phosphorylate the SMAD1/5, P38, and ERK1/2 MAPKs during the ameloblastic differentiation of iPS cells, only inhibition of SMAD1/5 phosphorylation by the inhibitor of ALK receptors, could reverse the ameloblastic differentiation of iPS cells. Taken together, the results showed that the ameloblastic differentiation of iPS cells induced by ameloblast-conditioned medium is a SMAD1/5-dependent, but MAPKs-independent, transcriptional event.

The results showed that iPS cells stimulated by ameloblast-conditioned medium for 14 days exhibited higher gene and protein levels of ameloblast markers AMBN, ENAM, and CK14, but a very low level of odontoblast markers DMP-1 and DSP, which agreed with previous results. These results suggested that cultures using conditioned medium from ameloblasts could induce iPS cells to differentiate into cell types that closely resemble ameloblasts.

BMPR-II plays a key role in the BMP4 signaling pathway. Previous studies have suggested that the BMP4 signal regulates the formation of Hertwig’s epithelial root sheath and the epithelial-mesenchymal interactions that occur during the patterning of the tooth crown and root. Meanwhile, BMPR-II is expressed substantially in the epithelial and mesenchymal cells of tooth germs at different stages of development. Therefore, in this study, we investigated the role of BMPR-II in ameloblastic differentiation of iPS cells induced by ameloblast-conditioned medium. As expected, the mRNA and protein levels of BMPR-II increased during the differentiation of iPS cells into ameloblasts. More importantly, silencing of BMPR-II using a specific siRNA reversed significantly the increased gene and protein levels of ameloblast markers AMBN, ENAM, and CK14 induced by the ameloblast-conditioned medium, confirming

**p < 0.01, vs. control group. **p < 0.01, vs. ASF-CM group.

Figure 5. RT-PCR analysis of the expression of Ambn, ENAM, and Ck14 of iPS cells induced by the control medium, ASF-CM, or ASF-CM respectively supplemented with 200 nM LDN-193189 (LDN, a selective SMAD1/5 inhibitor), 10 μM SB203580 (SB, a selective P38 inhibitor), or 10 μM U0126 (a selective ERK1/2 inhibitor).
that BMPR-II-mediated signal transduction plays a major role in the ameloblastic differentiation of iPS cells. Further study is needed to confirm the present phenomena by overexpressing BMPR-II transgene into iPS cells in the presence of ASF-CM. In addition, it was possible that factors present in ASF-CM could increase the expression of BMPR-II through Alk-Smad1/5 signal transduction, and this molecular mechanism needs to be further clarified.

The intracellular signal transduction of BMPR-II has been studied extensively. Although activation of SMAD1/5 is regarded as the canonical BMP...
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Smad1/5 signal transduction regulates the ameloblastic differentiation of induced pluripotent stem cells. The Western blotting results showed a significant increase in SMAD1/5, P38, and ERK1/2 phosphorylation after induction by ameloblast-conditioned medium. However, further analysis showed that only blockage of the SMAD1/5 pathway with the inhibitor of Alk receptors could decrease the gene and protein levels of AMBN, ENAM, and CK14. Neither the P38 pathway inhibitor SB203580 nor the ERK1/2 inhibitor U0126 has any reversing effect on the ameloblastic differentiation of iPS cells. Therefore, collectively, our data indicated that the ameloblastic differentiation of iPS cells induced by ameloblast-conditioned medium was regulated primarily by SMAD1/5 signal pathways activated by Alk receptors. However, this fact does not reduce the participation of BMPR-II in the ameloblast differentiation process because it has been reported that there is dimerization effect between Alk and BmpRII receptors.

The development and formation of teeth is very complicated and is regulated by many growth factors and signal pathways, and this dynamic process is unlikely to be manipulated only by the molecular factors identified to date. Therefore, in the present study, ameloblast-conditioned medium was used to induce ameloblastic differentiation of iPS cells. In addition, it has been reported that the P38 MAPK pathway is required for the effects of BMP2 and BMP7 on tooth development, such as the expression of p21 in the embryonic enamel knot and the expression of enamelin, ameloblastin, and β4-integrin in ameloblasts; mice with a deletion of P38 in the ectoderm display defective secretion of dental enamel and the absence of dental cusps. ERK 1/2 MAPK has been reported to play an important role in TGF-β1-induced growth, collagen turnover, and differentiation of stem cells from tooth apical papilla. Therefore, further study is needed to identify the involvement of the P38 and ERK1/2 pathways in the regulation of ameloblastic differentiation of iPS cells induced by growth factors within the ameloblast-conditioned medium, such as BMP2, BMP7, and TGF-β1.

There are some limitations in the present study. Firstly, only one murine iPS cell line was used, so the present results need to be confirmed by other iPS cell models. Secondly, although it has been shown that the ameloblast-conditioned medium contains several growth factors including BMPs, TGF-βs, Notch-1, and fibroblast growth factors, the present study did not clarify the active components within the ASF-CM. In order to assure the reproducibility of the experiments, we have mixed together all the supernatants during ameloblasts culture, and then used them as ASF-CM for the subsequent experiments. In this way, the active components in the conditioned media were thought to be the same among the experiments. Further ELISA assays are needed to provide more details of the active components within the ASF-CM used in the present study. Thirdly, it is important to confirm the results using stably transfected clones of siRNA. However, the screening of such clones might affect the biological behaviors of iPS cells, so in the present study we only used the transient transfection, and the Western blot results showed that the protein levels of BMPR-II have been significantly decreased after 24 h transient transfection of BMPR-II siRNA. Furthermore, the transient transfection of BMPR-II siRNA significantly reversed the increased expression of ameloblastin, enamelin, and cytokeratin-14 induced by ASF-CM.

In summary, we have demonstrated that the smad1/5 signaling transduction, activated by Alk receptors, regulates the ameloblastic differentiation of iPS cells induced by ameloblast-conditioned medium. Further in vivo experimental studies are required to confirm the effect of the SMAD1/5 signal transduction on the ameloblastic differentiation of iPS cells.

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