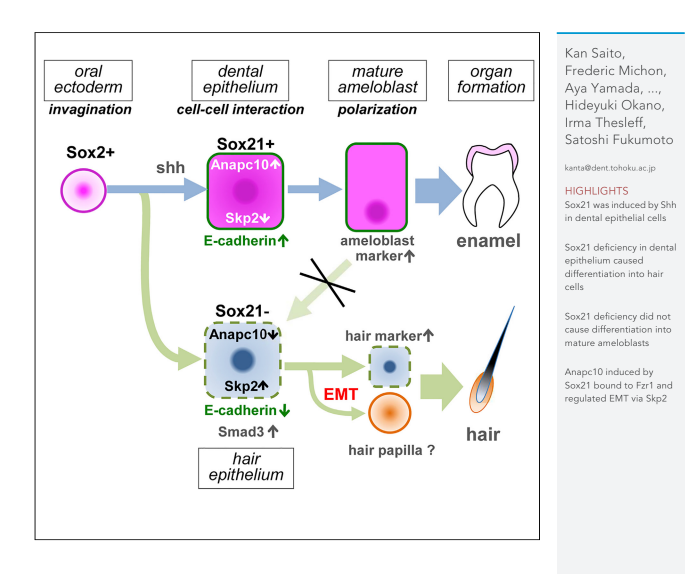
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Sox21 Regulates Anapc10 Expression and Determines the Fate of Ectodermal Organ



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SUMMARY

The transcription factor Sox21 is expressed in the epithelium of developing teeth. The present study aimed to determine the role of Sox21 in tooth development. We found that disruption of Sox21 caused severe enamel hypoplasia, regional osteoporosis, and ectopic hair formation in the gingiva in Sox21 knockout incisors. Differentiation markers were lost in ameloblasts, which formed hair follicles expressing hair keratins. Molecular analysis and chromatin immunoprecipitation sequencing indicated that Sox21 regulated Anapc10, which recognizes substrates for ubiquitination-mediated degradation, and determined dental-epithelial versus hair follicle cell fate. Disruption of either Sox21 or Anapc10 induced Smad3 expression, accelerated TGF- β 1-induced promotion of epithelial-to-mesenchymal transition (EMT), and resulted in E-cadherin degradation via Skp2. We conclude that Sox21 disruption in the dental epithelium leads to the formation of a unique microenvironment promoting hair formation and that Sox21 controls dental epithelial differentiation and enamel formation by inhibiting EMT via Anapc10.

INTRODUCTION

Members of the SRY-Box (Sox) B group share a conserved eight-amino-acid group B homology domain located adjacent to the HMG domain. The SoxB1 (Sox1–3) proteins function primarily as activators, whereas the closely related SoxB2 proteins (Sox14 and Sox21) are transcriptional repressors (Argenton et al., 2004; Sandberg et al., 2005). Furthermore, Sox21 was identified through proteomics studies to be associated with Sox2 during embryonic stem cell differentiation (Mallanna et al., 2010). The transcriptional activation/repression balance is important during the development of specific tissues. For example, SoxB2 repression of SoxB1 expression was shown to promote neural differentiation in the central nervous system (CNS) and peripheral nervous system (Sandberg et al., 2005; Uchikawa et al., 1999). This repression is supported by the expression of Sox21 throughout the developing CNS and brain (Cunningham et al., 2008). In addition, a major role of Sox21 has been demonstrated during hair shaft cuticle differentiation (Kiso et al., 2009) and its deletion affects the hair lipid composition (Kawaminami et al., 2012). However, the SoxB1 group proteins and their roles have received greater attention to date (Donner et al., 2007; Driskell et al., 2009; Groves and Bronner-Fraser, 2000) than SoxB2 group involvement in developmental processes.

The development of most ectodermal organs is initiated from epithelial thickenings called placodes, and their morphogenesis involves invagination and folding of the epithelium regulated by reciprocal interactions between the mesenchyme and epithelium (Dhouailly, 2009). The cross talk between both tissues involves specific molecular signals, such as Wnt, bone morphogenetic protein (BMP), sonic hedgehog (Shh), Fgf, Eda, and Tgf (Jernvall and Thesleff, 2012; Liu et al., 2016; Miyazaki et al., 2016). The process of ectodermal organ morphogenesis is highly conserved and largely regulated by the same genes, hence various developmental defects are often observed concordantly in several ectodermal organs. For example, patients with syndromes such as incontinentia pigmenti (Smahi et al., 2000), Langer-Giedion (Momeni et al., 2000), Ellis-van Creveld (Ruiz-Perez et al., 2003), tricho-dento-osseous (Price et al., 1998), anhidrotic ectodermal dysplasia (Srivastava et al., 1996; van der Hout et al., 2008), hidrotic ectodermal dysplasia (Han et al., 2018; Lamartine et al., 2000), Hallermann-Streiff (Pizzuti et al., 2004), and Menkes (Tumer et al., 2003) have dysplasia in both teeth and hair.

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The continuously growing rodent incisor represents a useful model to study stem cell regulation and organ development. Dental epithelial stem cells are localized in the proximal end of the incisor, and they express Sox2 and the Wnt inhibitor, Sfrp5 (Juuri et al., 2012). Dental epithelial cells differentiate into four types of epithelia: inner enamel epithelium (EE) and outer EE, stratum intermedium, and stellate reticulum. Inner EE expresses Shh, complementarily to Sfrp5, and differentiates into enamel-forming ameloblasts that express enamel matrix proteins, including amelogenin (Amel), enamelin (Enam), and ameloblastin (Ambn). Disruption of Amel or Ambn led to severe enamel hypoplasia, whereas hair abnormalities were not observed (Fukumoto et al., 2004; Gibson et al., 2001), indicating that these enamel matrix molecules are important for dental epithelium differentiation and enamel formation but not for hair development. Ameloblastin is critical for ameloblast differentiation in induced pluripotent stem cell-induced dental epithelium (Arakaki et al., 2012). In hair, the invaginated skin epithelium differentiates into interfollicular epidermis and hair follicles. After birth, adult stem cells residing in the basal layer of the epidermis and in the hair follicle bulge continuously regenerate the epidermis and hair follicles. Hair follicle stem cells derive from the bulge and migrate from the outer to the inner root sheath, where they express Keratin (Krt) 1, Krt10, Krt15, and Krt23 as epidermal keratins (Jensen et al., 1991; Rogers et al., 2004), as well as Krt27 and Krt32 as hair keratins (Langbein et al., 2010).

The present study focused on the role of Sox21 in tooth development. Although deletion of Sox21 is known to induce hair defects in mice (Kiso et al., 2009), deletion of the chromosome region 13q (containing the *SOX21* gene) in humans leads to irregular/dysplastic teeth (Kirchhoff et al., 2009).

RESULTS

Sox21 Is an Ameloblast Marker Regulated by Shh

The expression of *Sox21* mRNA during the tooth differentiation process was examined using *in situ* hybridization (Figure 1A). On embryonic day 15 (E15), *Sox21* was not detected in the dental tissue, but rather in part of the lip epithelium and in the whiskers. From E16 onward, *Sox21* expression was found in differentiating ameloblasts on the labial side of the incisor. *Sox21* expression was reinforced during ameloblast differentiation and at postnatal day 2 (P2); strong *Sox21* expression was detected in differentiated ameloblasts in the incisor and the molar (Figure 1A). To validate our findings, we used a reporter mouse carrying a GFP knockin at the *Sox21* locus (Kiso et al., 2009). As expected, GFP was detected in areas harboring differentiated ameloblasts in the mouse incisor and molar, i.e., the incisor labial side and the molar crown, respectively (Figure 1A). To evaluate the dynamics and specificity of *Sox21* expression, we monitored the expression by qPCR during tooth development and postnatal maturation. Although *Sox21* was not expressed at E13 and E14, its expression gradually increased between E15 and P7, reflecting the cell commitment and differentiation during tooth morphogenesis (Figure 1B). Furthermore, only epithelial cells isolated from the P1 incisor expressed *Sox21*, similar to the differential expression in the SF2 ameloblast cell line when compared with a mouse mesenchymal dental pulp cell line (Figure 1B).

As previously reported, Shh represents a marker of ameloblast differentiation (lseki et al., 1996). Therefore, we analyzed *Shh* and *Sox21* expression in mutants exhibiting abnormal ameloblast differentiation. As expected, the modification of the region of ameloblast differentiation was reflected through *Shh* and *Sox21* expression. Follistatin is a BMP antagonist that regulates ameloblast differentiation. We used *Follistatin* knockout (KO) and *K14-Follistatin* transgenic mice to confirm whether Sox21 is a marker of ameloblasts. Specifically, ectopic ameloblast differentiation in the lingual side of the incisor, as observed in *Follistatin* KO mice (Saville, 1970), was correlated with the expansion of *Shh* and *Sox21* (Figure 1C). Similarly, the loss of ameloblast differentiation in *K14-Follistatin* mice (Wang et al., 2004) was associated with a lack of *Shh* and *Sox21* expression in the labial side (Figure 1C). Taken together, our results indicated that *Sox21* specifically marks the ameloblast lineage during terminal cell differentiation.

Ameloblasts are constantly renewed in the ever-growing mouse incisor. Their maturation begins at the cervical loop (CL), which is located posteriorly, and continues while the cells migrate anteriorly in the EE. To investigate the possible association between *Sox21* expression and Shh signaling, we first investigated the expression patterns of Shh signaling pathway genes in the mouse incisor (Figure S1A). Although *Sox21* expression was initiated only in differentiated ameloblasts, the Shh pathway genes *Shh*, *Gli1*, *Gli2*, and *Ptc2* were expressed in the CL and early EE. These results indicate that Shh activity preceded *Sox21* expression further, we cultured E16 mouse incisors with or without the Shh inhibitor, cyclopamine. Through *in situ* hybridization, we showed that the absence of Shh activity greatly reduced the area of *Sox21* expression (Figure S1B). We confirmed this result by qPCR, which

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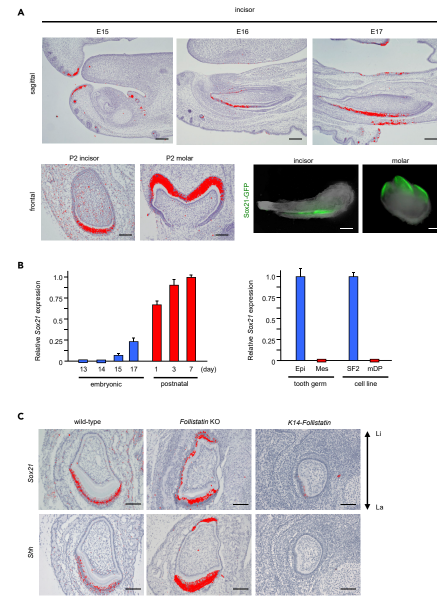


Figure 1. Expression of Sox21 in the Development of Incisor and Molar Processes

(A) Radioactive *in situ* hybridization (RISH) to analyze Sox21 expression in mice at embryonic day (E) 15, 16, and 17 and postnatal day (P) 2. Upper panels, sagittal sections of incisors; lower left panels, frontal sections of incisor and molar; lower right panels, GFP expression in dissected incisors and molars; fluorescence is from Sox21-GFP in Sox21+/- P2 mice. Scale bar, 200 μ m.

(B) Sox21 expression analyzed by qPCR. Tooth germs of each stage from E13 to P7 were collected (left panel). Dental epithelial cells and dental mesenchyme cells were isolated from the P1 tooth germ. In addition, SF2 cells (ameloblast cell line) and mDP cells (dental pulp cell line) were used for qPCR (right panel). Error bars represent mean \pm SEM of five technical replicates.

(C) In situ hybridization for the expression of Sox21 and Shh in mutant mouse incisors. Scale bar, 200 μ m. Li, lingual side; La, labial aspect of mandibular tooth.

demonstrated that recombinant Shh enhances *Sox21* expression (Figure S1C) as rapidly as 4 h after initiation of treatment. The addition of recombinant Shh in the culture medium enhanced *Sox21* expression, whereas the presence of cyclopamine drastically reduced *Sox21* expression in the E16 incisor and P2 dental epithelium (Figure S1C). Based on the rapid effect of Shh activity on the expression of *Sox21*, we suggest that the Shh pathway directly regulates *Sox21* expression during ameloblast differentiation.





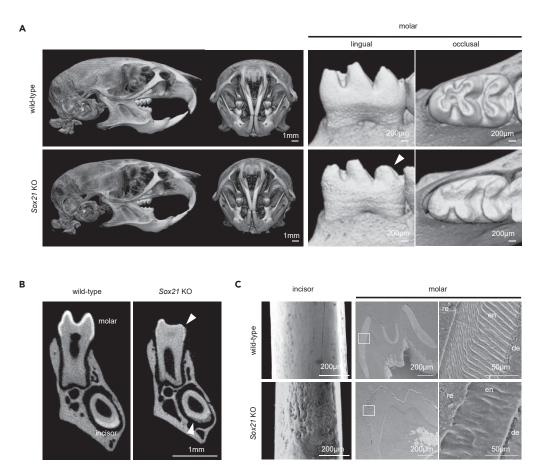


Figure 2. Phenotype of the Hard Tissue in Sox21 KO Mouse

(A–C) Wild-type and Sox21 KO mice (6-week-old) as analyzed by micro-CT (A and B) and SEM (C) for tooth formation. (A) Stereo images of the whole head, skull, and tooth (molar). Arrowhead indicates the excessive dental attrition. (B) Tomogram of left mandibular tooth. Arrowhead indicates the enamel of the Sox21 KO mouse. (C) SEM image showing tooth surface of the incisor (left panel). Low (middle panel) and high (right panel) magnification of a molar cross-section obtained by SEM. de, dentin; en, enamel; re, epoxy resin.

Sox21-Deficient Mice Display Signs of Amelogenesis Imperfecta

To understand the role of Sox21 expression during ameloblast differentiation, we assessed the teeth of Sox21 KO mice at 6 weeks of age, using micro-computed tomography (CT). In the head region, the size of the entire head remained unchanged and the molars exhibited excessive dental attrition in Sox21 KO mice (Figure 2A). No morphological changes were observed in the dentin and roots of molars, but the surface of the enamel was seriously affected and rugged in both molars and incisors (Figures 2A and 2B). The enamel thickness in the molar crowns was markedly reduced (Figure 2B arrowhead). Scanning electron microscopy (SEM) showed a seriously rugged surface of the enamel in the incisors and the loss of enamel prism structure in the molars (Figure 2C). The evaluation of tooth calcification by micro-CT further indicated a drastic reduction in enamel density. Sox21 KO incisors and molars showed only 9% and 6% of the enamel volume of the controls (Figure S2), although the dentin layer remained unaffected. Thus, we concluded that Sox21 is instrumental for proper enamel layer deposition.

Sox21 Deficiency Leads to Loss of Ameloblast Polarity and a Change in Cell Fate toward Skin Identity

As Sox21 is expressed in ameloblasts and loss of Sox21 leads to enamel defects, we analyzed the morphology of the ameloblast layer in 6-week-old Sox21 KO mice. In wild-type mice, the ameloblasts formed a continuous epithelial monolayer of elongated polarized cells. The nuclei were at the cell base, and the enamel layer was deposited from the cell apex (Figure 3A). Upon Sox21 deficiency, the ameloblasts were poorly elongated, and some cells had lost their polarity. Some cells were embedded in the enamel layer, forming pits within the

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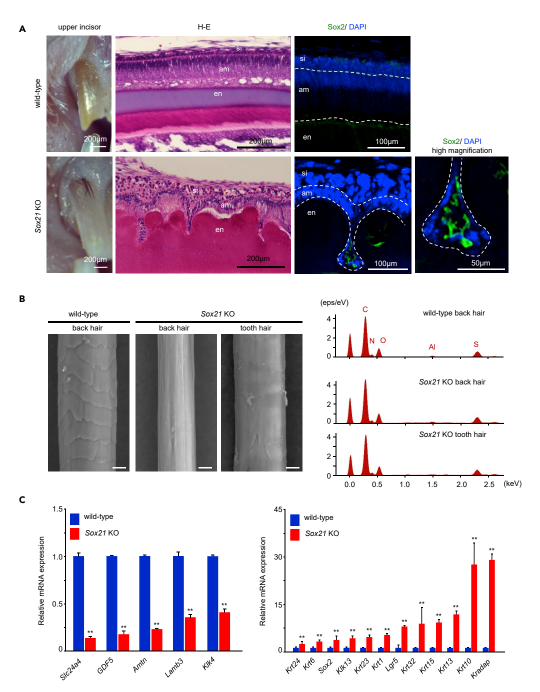


Figure 3. Histological Analysis of the Sox21 KO Mouse Incisor

(A) The upper incisors of 5-week-old wild-type and Sox21 KO mice were observed using a stereo microscope (left panel). The lower incisors of a wild-type and a Sox21 KO mouse were stained with hematoxylin and eosin (H&E) (middle panel). The right panels show high magnifications of ameloblasts in an incisor. The ameloblasts of the mouse incisor were immunostained using anti-Sox2 antibody with DAPI (right panel). si, stratum intermedium; am, ameloblasts; en, enamel.
(B) Surface structure of a wild-type back hair, Sox21 KO back hair, and Sox21 KO tooth hair as imaged by SEM. Scale bar, 10 μm. The elemental mapping of the hair was analyzed by energy-dispersive X-ray spectrometry.

(C) qPCR analysis of the expression of genes identified as being repressed or induced in the microarray data. P1 tooth germs of wild-type and Sox21 KO mice were used for qPCR. Error bars represent mean \pm SEM of five technical replicates. Student's t test (**p < 0.001).



mineralized tissue (Figure 3A). Furthermore, the cells at the bottom of the pits expressed Sox2. Sox2 expression has been shown to mark dental epithelial stem cells in the cervical loop of the mouse incisor (Juuri et al., 2012); Sox2 is then markedly downregulated in differentiated ameloblasts (Figure S3). Cell proliferation was analyzed by 5-ethynyl-2'-deoxyuridine (EdU) staining. EdU assays measure the incorporation of EdU into newly synthesized DNA. Sox2 expression remained in the differentiated dental epithelium, such as ameloblasts (Figure S3 arrowhead), whereas expression of Sox2 did not affect cell proliferation because EdU was unchanged in *Sox21* KO mice (Figure S3). In the embryonic stage of incisor and molar development, no histological differences in tooth germ layers were observed between wild-type and *Sox21* KO mice (Figure S4A). Furthermore, the disorganization of the dental epithelium was observed on postnatal day 7 molars in *Sox21* KO mice (Figure S4B arrow). In the case of incisors, the dental epithelium from the embryonic stage to postnatal day 7 was normal (Figure S4C). However, disorganization and lost polarity were obtained in the anterior dental epithelium of P4-week-old incisors, and the epithelium invaginated and formed pits at 6 week (Figure S4D and S4E). Therefore, the expression of *Sox2* in the bottom of the enamel pits reflected dedifferentiation of the ameloblasts to remaining Sox2-positive dental epithelial stem cells.

Oral examination revealed that in 63% of 5- to 6-week-old Sox21-deficient mice, hairs appeared between the tooth surface and the gingival sulcus in both the lower and upper jaws (Figures 3A, 5A, and S5A), but not in the molars. Sox21 was not expressed in the early embryonic stage (Figures 1A and 1B), and the ameloblasts had normal morphology until P7 in Sox21 KO mice incisors (Figures S4A and S4C). The cell arrangement of the ameloblast layer was disturbed in postnatal 4-week-old incisors (Figure S4D). However, ectopic hair was not formed in this stage. Hair was formed within the gingiva at 5 weeks and erupted at 5 to 6 weeks (Figure S5A). From this result, ectopic hair was formed in the secretory stage of the ameloblast layer at around 5 weeks, but not the early stage of tooth development. Ectopic hair contains medulla and cortex structure of the hair, as well as follicle papilla cells (Figure S5B). Such incomplete hair is observed with teratomas and dermoid cysts (Ahmad et al., 2013; Komiyama et al., 2002; Lewis et al., 2012; Takeda et al., 2003; Tavares et al., 2018). However, no typical bulge structure was observed, and we did not find any teratomalike structure in Sox21 KO incisors. Previously, it was reported that Sox21 deficiency leads to a modification of the hair cuticle layers, resulting in defective anchorage, and. ultimately, to cyclic alopecia (Kiso et al., 2009). Thus, we compared the surfaces of dental and back skin hairs. The surface of the back hairs was smooth in Sox21 KO mice and different from that of the wild-type mice (Figure 3B). Notably, the tooth hairs in Sox21 KO mice displayed a smooth surface, reflecting the lack of a cuticle, which was similar to the back hair in Sox21 KO mice. Furthermore, analysis of the elemental mapping at the compositional and microstructural levels by energy-dispersive X-ray spectrometry revealed high similarity between the different hair types (Figure 3B). From macroscopic analyses, some back hairs appeared thin with zigzag shape, whereas the tooth hairs were short and straight (Figure S5C). The tooth hairs were thicker than the back hairs (Figures 3B and S5C). Therefore, the length of each hair was measured (Figure S5D). The average length of the tooth hairs was 2.49 \pm 0.36 mm, whereas that of the back hairs was 3.19 \pm 0.32 mm (p < 0.005, Student's t test). Tooth hairs were significantly shorter than back hairs.

To investigate whether ectopic hairs of *Sox21* KO mice switched tooth cell fate, microarray analysis were performed between tooth and skin in wild-type (Table S1). Conversely, almost all the genes upregulated in *Sox21* KO teeth were more highly expressed in the skin. Notably, numerous keratin-related genes were upregulated in the *Sox21* KO tooth germ and wild-type skin. The microarray data were validated by qPCR. Specifically, *Amtn* and *Klk4* were decreased in *Sox21* KO ameloblasts (Figure 3C), whereas the expression of *Shh*, *Gli1*, *Gli2*, *Ptc2*, *Mmp20*, and *Amel* remained unchanged (data not shown). As for the microarray results, the expression of dental epithelial stem cell marker *Sox2*, as well as keratin family and hair-related genes, such as *Lgr5*, were increased in *Sox21* KO dental epithelium (Figure 3C). Lgr5 is expressed in hair germ and hair bulge of the hair root sheath wall region (Leung et al., 2018), whereas Sox2 is expressed in the dermal papilla of the hair root sheath bottom (Driskell et al., 2009). The expression of Sox2 and Lgr5 was examined by immunostaining in 6-week-old mouse incisors (Figures 3A and S6A). These expression patterns were similar to that in the hair root sheath, as Sox2 was expressed in the bottom and Lgr5 was expressed in the wall of the ameloblast invagination pit. Taken together, these results reflected the perturbed differentiation of ameloblasts and a cell fate change from dental to skin identity in the *Sox21* KO incisors.

Anapc10 Is a Sox21 Direct Target Gene Involved in Ameloblast Differentiation

To determine the role of Sox21 in amelogenesis, we attempted to determine Sox21 direct target genes via chromatin immunoprecipitation sequencing analysis (Figure 4A). Among the direct targets identified, a

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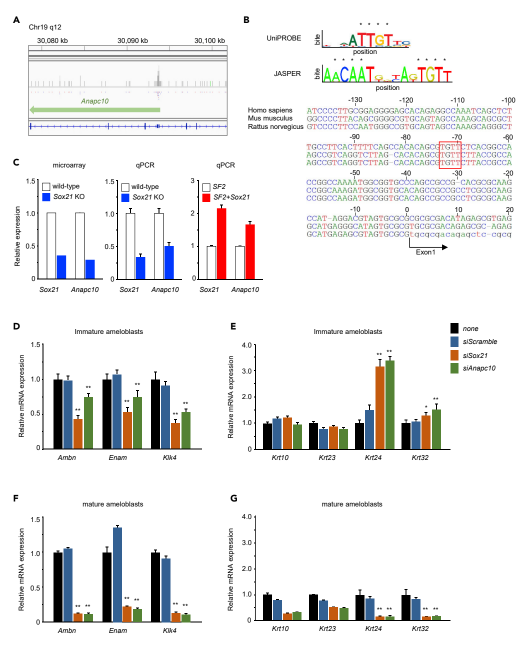


Figure 4. Identification of Sox21 Downstream Molecules and Its Effect on Ameloblasts

(A) Chromatin immunoprecipitation sequencing analysis of Sox21 interactors in SF2 cells. The peak of bound chromatin fragments representing *Anapc10* was visualized using IGV software.

(B) Sox21 binding sites were investigated in UniPROBE and JASPAR databases (top). The upstream of Sox21 of Homo sapiens, Mus musculus, and Rattus norvegicus were aligned. Capital letters represent intronic nucleotides. (C) Expression levels of Sox21 and Anapc10 in the ameloblasts of Sox21 KO and wild-type mice as investigated by microarray analysis (left) and qPCR (middle). Sox21 overexpression in SF2 cells as examined by qPCR (right). Error bars represent mean \pm SEM of five technical replicates.

(D–G) Expression of Sox21 and Anapc10 in SF2 cells was inhibited by siRNA. SF2 was cultured by NT-4 for 48 h to differentiate to mature ameloblasts (F and G). Scramble siRNA was used as a negative control. After 48 h, the expression of ameloblast markers (D and F) and keratin family genes (E and G) was evaluated by qPCR. Error bars represent mean \pm SEM of five technical replicates. Student's t test (*p < 0.005, **p < 0.001).





Sox21 binding peak was detected in the promoter region of anaphase-promoting complex subunit 10 (Anapc10) (Figure 4A). Subsequently, a candidate Sox21-binding consensus sequence (Matsuda et al., 2012) was identified in the Anapc10 promoter region, using the UniPROBE (http://thebrain.bwh.harvard. edu/uniprobe/) and JASPER (http://jaspar.genereg.net) databases (Figure 4B). qPCR validation of the effects of Sox21 deficiency or excess on the expression of these target genes indicated that only Anapc10 expression was modulated by the Sox21 level (Figure 4C). These results indicate that the expression of Anapc10 was dependent on Sox21 in ameloblasts. Anapc10 has previously been shown to regulate the cell cycle through cyclin B ubiquitination (Kominami et al., 1998). However, the role of Anapc10 in the dental context was unclear. Therefore, we examined the effects of Sox21 and Anapc10 on the expression of Ambn, Enam, and Klk4, which are markers of proach to detect molecular changes induced by Sox21 deficiency (Table S1). We dissected tooth germs from P1 Sox21 KO and control incisors. In particular, several markers of ameloblast maturity (i.e., amelotin [Amtn], laminin α 3 [Lama3], and kallikrein-related peptidase 4 [Klk4]) were included among the downregulated molecules in the Sox21 KO tooth germ. Hair and tooth development-related genes were largely similar between the P1 incisor tooth germ, including epithelium and mesenchyme, and P1 skin, including hair follicles (Thesleff and Mikkola, 2002). Except for Wnt16, genes that were downregulated in the Sox21 KO tooth germ were also differentiated ameloblasts in the rat dental epithelial cell line SF2. SF2 is one of the immature (undifferentiated) dental epithelial cell lines that differentiates into ameloblasts when stimulated with neurotrophin 4 (NT-4) in vitro (Yoshizaki et al., 2008). Upon knocking down either Sox21 or Anapc10, immature and mature ameloblasts reduced ameloblast marker expression (Figures 4D and 4F). Furthermore, the expression of dermal keratin Krt24 and hair keratin Krt32 was increased following Sox21 or Anapc10 knockdown in immature ameloblasts (Figure 4E). In differentiated SF2 cells, after stimulation by NT-4 and knocking down either Sox21 or Anapc10, no hair marker expression was observed (Figure 4G). Our results suggest a genetic network involving Sox21 and Anapc10 upstream of Ambn, Enam, and Klk4 during the terminal differentiation of ameloblasts. Furthermore, immature ameloblasts (such as pre-ameloblasts), but not differentiated ameloblasts, may undergo trans-differentiation into hair cells from dental epithelium.

Sox21 Disruption Enhances Epithelial-To-Mesenchymal Transition and TGF- $\beta 1$ Signaling in Dental Epithelium

We next examined the cells forming the ectopic tooth hairs. The histological analysis of the tooth hair showed presence of a hair shaft and a population of cells at the hair root (Figures 5A and S5B). As the hair follicle structure was not clear on sagittal sections, we used frontal sections to visualize this structure. Notably, we detected Sox2 expression in the hair root sheath localized in the enamel pits, whereas amelo-blastin was absent from the hair structures (Figure 5A).

In the tooth hair, morphogenesis relies on cross talk between the epithelial and mesenchymal compartments. The development of ectopic tooth hairs upon Sox21 deficiency implies the presence of a dermal papilla at the base of the hair. To investigate this, we analyzed the presence of dermal markers in the bottom of the enamel pits. Expression of Sox2, which is known to be a dermal papilla marker (Driskell et al., 2009), was detected in the pits (Figures 3A and 5A). To examine this further, we assessed the presence of additional dermal papilla markers, N-cadherin and Vimentin. Although these markers were only weakly expressed in differentiated ameloblasts, loss of Sox21 induced strong expression of both cadherins in the ameloblast layer (Figure 5B). Furthermore, N-cadherin was strongly expressed in the root sheath of the Sox21 KO tooth hairs. Similarly, pan-keratin was detected in enamel matrix-secreting mature ameloblasts and in the papillary layer, and Sox21 deficiency induced strong N-cadherin and Vimentin expression in the tooth hair root sheath (Figures 5B and S6B arrow). As our results suggested the appearance of dermal cells at the base of the tooth hair, we subsequently investigated the impact of Sox21 and Anapc10 knockdown on mesenchyme markers, such as N-cadherin, Vimentin, Zeb1, Zeb2, and Snail1, in the SF2 cell line (Figures S7A and S7B). As anticipated, the mesenchymal markers were all upregulated upon Sox21 or Anapc10 silencing in only immature ameloblasts. Our results therefore support the roles of Sox21 and Anapc10 in the maintenance of epithelial identity; when one of these are lacking, the differentiating ameloblasts can initiate epithelial-to-mesenchymal transition (EMT).

EMT is primarily induced through Smad3 and 4, which are under the control of Tgf activity (Kaimori et al., 2007; Zavadil et al., 2004). Therefore, we used small interfering RNAs (siRNAs) in SF2 cells to suppress the levels of Sox21 and Anapc10 and analyze the impact on Smad3, Smad6, and Smad7 by western blotting. The silencing of Sox21 and Anapc10 did not affect the Smad 6 and 7 levels in the cells; however, the amount

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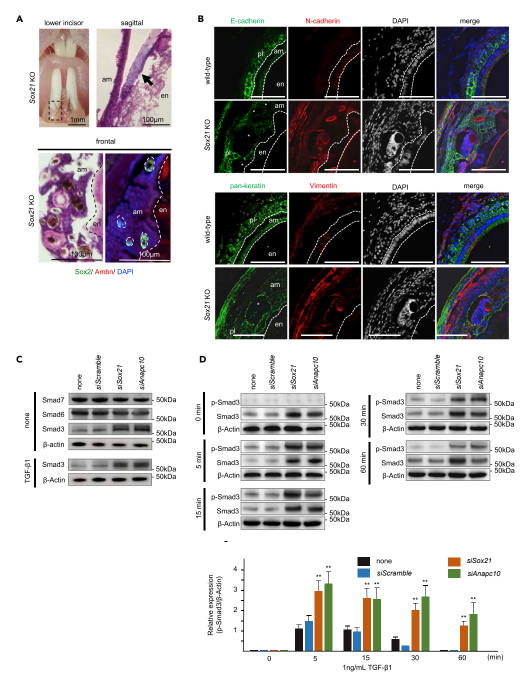


Figure 5. Molecular Analysis of the Ameloblasts in the Sox21 KO Mouse

(A) Histological analysis of the *Sox21* KO mouse lower incisor gingiva. H&E staining of sagittal and frontal sections of the hair shaft in ameloblasts and immunostaining of Sox2 and Ambn in the frontal section. am, ameloblast; en, enamel. (B) Double immunostaining of incisal frontal section from a 6-week-old mouse, counterstained with DAPI. Upper panels show sections stained using anti-E-cadherin and anti-N-cadherin antibodies; for lower panels, we utilized anti-pan-keratin and anti-Vimentin. Scale bar, 200 μ m. am, ameloblast; en, enamel; pl, papillary layer. *root sheath analog. (C and D) Expression of *Sox21* and *Anapc10* in SF2 cells was inhibited by siRNA for 48 h (C) Smad3, 6, and 7 protein levels as detected by western blotting after SF2 cells were cultured with or without TGF- β 1 (1 ng/mL) for 24 h. (D) *Sox21*- or *Anapc10*-silenced SF2 cells were stimulated with TGF- β 1. After 5, 15, 30, or 60 min, Smad3 and p-Smad3 expression was analyzed by western blotting. The p-Smad3 expression was normalized to that of β -actin. The amount of p-Smad3 in control SF2 cells induced by TGF- β 1 after 5 min was calculated as a standard. Relative expression of p-Smad3 is indicated. Error bars represent mean \pm SEM of three technical replicates. Student's t test (**p < 0.001).





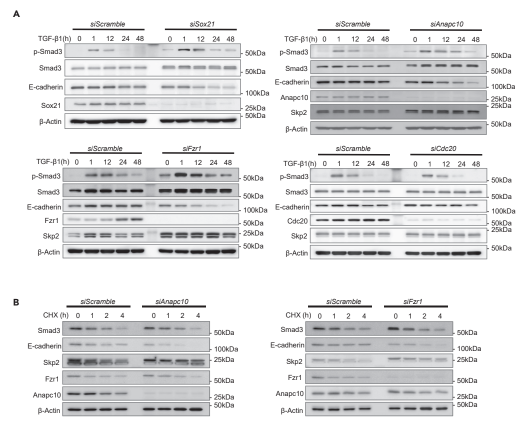


Figure 6. Regulation of EMT in SF2 Cells by Knockdown of Anapc10, Fzr1, and Cdc20

(A) SF2 cells were transfected with Sox21, Anapc10, Cdc20, or Fzr1 siRNA probes. After 48 h, cells were cultured with TGF- β 1 from 0 to 48 h and then processed for immunostaining.

(B) siRNA-transfected cells were treated with 20 μ g/mL cycloheximide (CHX) to determine the half-life of the indicated proteins. The cells were collected at the indicated time points for western blotting.

of Smad3 increased drastically (Figure 5C). Notably, these results were transforming growth factor (TGF)- β independent, because the increase in Smad3 did not change with or without TGF- β 1. To monitor the effects of *Sox21* and *Anapc10* silencing on TGF- β activity, we measured the kinetics of Smad3 phosphorylation upon TGF- β activation (Figure 5D). The effect of TGF- β 1 on the SF2 cells, visualized by the presence of phospho-Smad3, was greatly enhanced by *Sox21* and *Anapc10* knockdown as rapidly as 5 min after TGF- β 1 addition. These results suggest that the EMT process seen in *Sox21* KO ameloblasts originated from TGF- β activity through the phosphorylation of Smad3.

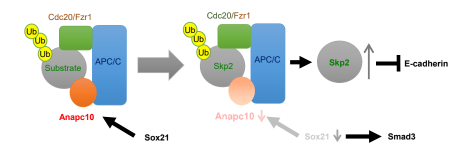
Anapc10 and Fzr1 Regulate EMT by Modulation of the Expression of EMT-Related Proteins

Anapc10 is a core component of the APC/C multi-subunit E3 ligase complex and is implicated in substrate recruitment to the complex by functioning as a substrate recognition module together with a co-activator protein, cell division cycle 20 (Cdc20) or CDC20 homolog 1 (Cdh1/Fzr1) (da Fonseca et al., 2011; Kominami et al., 1998). Thus, to investigate the contribution of APC/C activity to the acquired EMT phenotypes elicited by Sox21 deficiency in mice, we conducted an immunoblot analysis to analyze the abundance of the EMT-related proteins Smad3 and E-cadherin in SF2 cells upon knockdown of *Sox21*, *Fzr1*, *Cdc20*, or *Anapc10*. Knockdown of *Sox21*, *Fzr1*, or *Anapc10*, but not *Cdc20*, caused increase in Smad3 protein and induced Smad3 phosphorylation by TGF- β (Figure 6A). Furthermore, we observed an increase in N-cadherin and a reduction in E-cadherin upon *Fzr1* or *Sox21* knockdown in the SF2 cells (Figure S7D).

In addition, we identified a canonical D-box motif (RxxLxxxxN), which could be targeted by APC/C, in the Smad3 protein, prompting us to test the possibility that APC/C directly regulates Smad3 protein stability through post-translational modification. Hence, we determined the half-lives of Smad3 protein in *Fzr1* or *Anapc10* knockdown and control cells. No significant changes in the half-life of Smad3 protein following









Fzr1 or *Anapc10* knockdown were detected (Figure 6B). This result implies that Smad3 was not subjected to APC/C-mediated ubiquitination and degradation in this circumstance. In comparison, Skp2, a reported APC/C^{Fzr1} substrate (Bashir et al., 2004; Wei et al., 2004), was accumulated and stabilized following the depletion of Fzr1 or Anapc10 in SF2 cells (Figure 6B). Furthermore, consistent with a previous report showing Skp2-mediated E-cadherin degradation (Inuzuka et al., 2012), the protein levels and half-lives of Skp2 and E-cadherin displayed inverse correlations upon *Fzr1* or *Anapc10* knockdown in SF2 cells (Figure 6B). These findings indicate that Fzr1 and Anapc10, both of which play a crucial role in APC/C substrate recognition, regulate EMT by modulating the abundance and activity of EMT-related proteins such as Smad3, Skp2, and E-cadherin (Figure 7).

Knockdown of *Sox21*, *Fzr1*, or *Anapc10* Promotes TGF-β-Induced E-Cadherin Downregulation

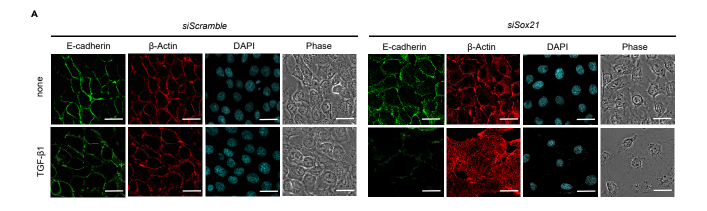
To further investigate the effects of Sox21 knockdown in the induction of EMT, we sought to analyze E-cadherin protein levels in Sox21, Fzr1, Cdc20, or Anapc10-depleted SF2 cells. Both E-cadherin and actin were largely confined to the cell membrane in control cells, as is normally observed in tooth epithelial cells in unstimulated conditions (McCrea et al., 1991) (Figure 8A). Even following TGF-β stimulation, E-cadherin and β -actin abundance on the cell membrane was not significantly altered, indicating that dental epithelial cells are not responsive to TGF-β stimulation under normal conditions (Figure 8). Conversely, knockdown of Sox21, Anapc10, or Fzr1, but not Cdc20, resulted in the downregulation of E-cadherin, which was further promoted upon TGF-β stimulation (Figures 8A, 8B, and S7E). Furthermore, we analyzed EMT marker expression in dental epithelial cells after transfection with Sox21 or Anapc10 siRNA cultured with TGFβ1 to induce EMT. EMT markers N-cadherin, Vimentin, Zeb1, Zeb2, and Snail1 were not induced by siRNA for Sox21 or Anapc10 in NT-4-induced differentiated ameloblasts (Figure S7B), but were induced in undifferentiated immature dental epithelium (Figure S7A). These experiments revealed that immature dental epithelium, but not differentiated ameloblasts, underwent trans-differentiation into hair cells and may induce EMT to form hair papilla. Furthermore, upon Sox21, Anapc10, or Fzr1 knockdown, β -actin was localized to the cell membranes in the absence of TGF- β 1, whereas it was abundantly localized throughout the cells in the presence of TGF-β1 (Figures 8A and 8C). Therefore, these data suggested that Sox21-induced Anapc10 expression represents a crucial element for suppressing EMT in dental epithelium, and that dysregulation of the inhibitory mechanism may cause an aberrant EMT, leading to abnormal formation of an ameloblast-derived root sheath analog.

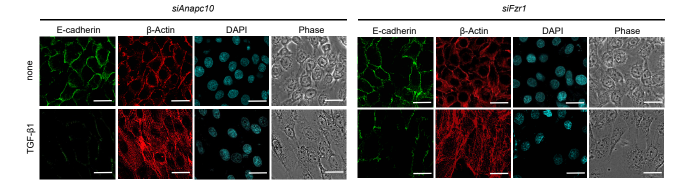
DISCUSSION

Our study revealed a role of Sox21 during ameloblast differentiation. We showed that Sox21 is expressed in ameloblasts, which differentiate from Sox2-expressing stem cells in dental epithelium (Juuri et al., 2012). Notably, Sox2 and Sox21 have been proposed to have a reciprocal relationship in the nervous system (Argenton et al., 2004; Sandberg et al., 2005). Sox2 expression in the incisor was observed in the cervical loop region and then decreased during dental epithelium differentiation in both wild-type and Sox21 KO mice (Figure S3). Furthermore, Sox2-positive dental epithelium re-appeared at a later stage in Sox21 KO incisors (Figure S3 arrowhead), indicating that Sox21 may inhibit Sox2 expression and form the ectopic hair organ. Some cells in the secretory stage of ameloblast start invagination and form a hair organ-like structure. Some pre-ameloblasts may have been retained as



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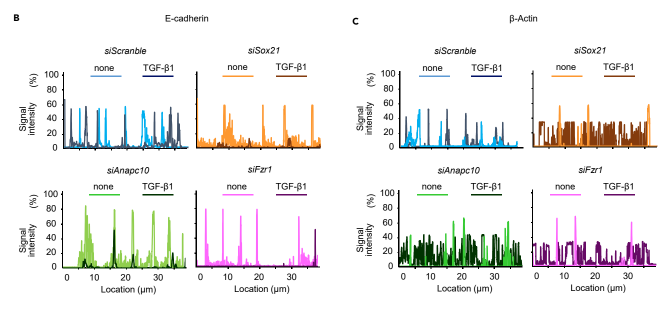


Figure 8. Abolishment of the Dental Epithelium Cytoskeleton by Sox21 and Anapc10 Suppression with TGF-β (A) SF2 cells in which Fzr1, Anapc10, or Sox21 was depleted by siRNA were cultured in the presence or absence of TGF-β1 for 48 h. Cells were stained with anti-Ecadherin antibody (green), anti-β-actin antibody (red), and DAPI (aqua). The protein localization was analyzed by confocal fluorescence microscopy. Scale bar, 10 µm. (B and C) Fluorescence intensity of E-cadherin (B) and β-actin (C) as evaluated with the ImageJ software (NIH).

ameloblast lineage cells, whereas others switched to hair-forming cells. EdU labeling in the incisor revealed that Sox2-positive cells observed in *Sox21* KO mice did not proliferate and may therefore not generate progeny (Figure S3). Expression of many keratins was increased in the P1 mouse dental

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epithelium (Figure 3C), which was histologically unchanged (Figures S4A and S4C). Similar results were obtained with the immature ameloblast cell line (Figure 4E). The preparation of genes to make hair may already start from this time. In the incisor, epithelial invasion of a hair-like structure appeared at P4 weeks and hair was formed at P6 weeks (Figures S4D, S4E, and S5A). Lgr5 is expressed in hair germs and the hair bulge that makes up the hair root sheath wall, whereas Sox2 is expressed in the dermal papilla located at the bottom. Immunohistochemistry of Lgr5 and Sox2 in the 6week-old Sox21 KO incisor showed a similar expression pattern to that in hair roots (Figures 3A and S6A). The expression of these molecules, which is absent in ameloblasts, indicates that the ameloblasts were transformed into hair-forming cells. Sox2-positive cells are presented in some pre-ameloblasts and stratum intermedium, which may form ectopic hairs in the tooth germ. In the molar, lose polarization was observed in P7-day-old Sox21 KO mice (Figure S4B). However, we did not observe ectopic hairs in the molar region. The molar started eruption at P2 weeks, and dental epithelial cells were lost after this eruption. From this situation, it is possible that the molars did not have hair structures because they did not have sufficient time to grow. Previously, we showed that the basement membrane component nephronectin and NKX2-3 inhibit Sox2 expression in the dental epithelium (Arai et al., 2017; Han et al., 2018). However, nephronectin was downregulated in differentiated ameloblasts. Both Sox21 and nephronectin may inhibit Sox2 expression in the dental epithelium. Because $Sox21^+$ and $Sox2^+$ cell populations are not in close proximity in the incisor epithelium, they do not interact directly. However, our results suggested that Shh activity occurs directly upstream of Sox21 expression in ameloblasts. By using the predictive algorithm JASPAR, we identified that the main transcription factors predicted to bind upstream of the Sox21-coding sequence were Znf263, Egr1, and Gli2, of which Gli2 is activated by the Shh pathway. It was reported that Hedgehog signaling acts upstream of Sox2 in cancer stem cells (Gopinath et al., 2013), and Sox18 is a target gene of hedgehog signaling in cervical carcinoma (Petrovic et al., 2015). Therefore, we hypothesized that Shh may regulate Sox21 expression via Gli2 activity. Sox2-expressing cells in tooth epithelium have been shown to be targets of Shh, based on their Gli1 expression (Li et al., 2015). Shh induced Sox21 expression, and inhibitor of Shh signaling inhibited Sox21 expression (Figures S1B and S1C). From this result, Shh may be an upstream molecule of Sox21. The deletion of Shh shows no hair formation and presence of abnormal follicular structures (St-Jacques et al., 1998). Shh is important for the invagination of the tooth and hair epithelium. The deletion of Shh signaling may inhibit initial tooth and hair formation. Previous reports showed that ectopic hairs were observed in Med1 and FAM38h KO incisors (Wang et al., 2016c; Yoshizaki et al., 2014). Med1 regulates Notch1 and Ca²⁺ channel expression and is involved in ectopic hair formation. However, unaltered expression of Med1, FAM38h, Notch1, and Ca²⁺ channel in Sox21 KO mice indicated that Sox21, Med1, and FAM38h have independent functions in ectopic hair formation. Furthermore, the hair follicle structure of ectopic hair was unclear in Med1 KO mice. The hair follicle structure in ectopic hairs of teratomas and dermoid cysts were also unclear (Ahmad et al., 2013; Komiyama et al., 2002; Lewis et al., 2012; Takeda et al., 2003; Tavares et al., 2018). Our results similarly found unclear hair follicle structures. Thus, the hair follicle with ectopic hair may be immature and obscure.

Sox21 deficiency impaired ameloblast function, leading to decreased enamel volume and defective structure of the remaining matrix. The expression of Amtn, Ambn and Enam, which are important for enamel matrix formation and calcification (Meredith et al., 2014), was decreased. Furthermore, we showed that Sox21 directly regulates the expression of Amtn and Klk4, which are involved in ameloblast maturation and enamel formation (Gasse et al., 2012; Kawasaki et al., 2014). Unexpectedly, when Sox21 expression was inhibited, ameloblasts lost their dental identity and trans-differentiated into hair cells. It is possible that, without a clear terminal differentiation signal, the immature dental epithelium retains the capacity to adopt a different cell fate. SF2 cells differentiated into ameloblasts by NT-4. We analyzed the expression of hair cell markers in both differentiated ameloblasts and immature dental epithelium cells, using SF2 cells after transfection with Sox21 siRNA. In immature dental epithelium, Sox21 siRNA inhibited the expression of ameloblast markers and induced expression of hair keratins Krt24 and Ker32 (Figures 4D and 4E). However, in NT-4-induced differentiated ameloblasts, Sox21 siRNA inhibited ameloblast marker expression and did not induce hair markers (Figures 4F and 4G), indicating that immature dental epithelium, but not differentiated ameloblasts, could differentiate into hair cells. Interestingly, loss of Sox21 seemed to favor the EMT process, and it is possible that this led to the formation of dermal papilla. The acquisition of migratory properties is a prerequisite for cancer invasion into surrounding tissues. The acquisition of migration requires a dramatic morphologic alteration, termed EMT, such as loss of epithelial characteristics



of cell polarity and cell adhesion (Thiery et al., 2009). We speculate that the epithelial characteristic was lost by lack of Sox21, because the dental epithelial arrangement was disrupted and cell polarity was lost after 4 weeks in the *Sox21* KO mice (Figure S4D). Therefore, we analyzed EMT marker expression in dental epithelial cells after transfection with *Sox21* siRNA. EMT markers, *N-cadherin, Vimentin, Zeb1, Zeb2*, and *Snail1*, were not induced by siRNA for *Sox21* in differentiated ameloblasts, but were induced in immature dental epithelium (Figures S7A and S7B). It was reported that various cell adhesion molecules, such as cadherin, contribute to tooth differentiation (Chiba et al., 2019; Saito et al., 2015; Verstraeten et al., 2013; Yamada et al., 2016). These experiments revealed that immature dental epithelium, but not differentiated ameloblasts, *trans*-differentiated into hair cells and may induce EMT to form hair papilla. However, further experiments are needed to demonstrate whether the mesenchymal cells of the dermal papilla actually originated from the immature epithelial ameloblasts via EMT in the *Sox21* KO teeth. It is also possible that Sox2 positivity in the forming dermal papilla mesenchyme was induced by the epithelium after it had adopted hair fate by *Sox21* deletion.

EMT not only is induced by TGF- β signaling but also results from Sox2 activity (Gao et al., 2015; Wang et al., 2016b) and BMP (Dudley et al., 1995; Rothhammer et al., 2005), canonical Wnt (Kemler et al., 2004), and Notch (Xie et al., 2012) signaling. In the tooth, EMT can reportedly occur by the cessation of BMP and Wnt signaling, leading to a switch from crown to root formation (Yang et al., 2013). However, in the present study, the epithelium did not differentiate to mesenchymal structures, but generated the epithelial cell network covering the root (epithelial cell rests of Malassez). In turn, TGF- β activity is important for tooth formation and cell differentiation (Vaahtokari et al., 1991). Specifically, TGF signaling in ameloblasts (Khan et al., 2013; Zhu et al., 2000) is important for enamel formation as its inhibition can lead to amelogenesis imperfecta (Huckert et al., 2015; Poche et al., 2012; Zhao et al., 2011). Thus, TGF activity does not normally induce EMT in the dental epithelium. This conclusion is supported by our result demonstrating the maintenance of E-cadherin expression in SF2 cells upon the addition of TGF-B1 (Figure 8A). However, TGF- β activity was enhanced by decreases in Sox21 and Anapc10, leading to EMT induction *in vitro*. Furthermore, signs of EMT, i.e., the expression of Vimentin, and N-cadherin, were observed in vivo upon Sox21 deficiency. Thus, our results suggest that the maintenance of ameloblast identity and their proper differentiation depend on Sox21 expression. However, EMT did not occur in NT-4-induced differentiated ameloblasts in vitro (Figures S7A and S7B), although it may have occurred in the immature dental epithelium of Sox21 KO mice.

Our findings demonstrate that, among the Sox21 direct targets, the expression of Anapc10 is crucial in the ameloblast differentiation process. Anapc10 is one of the core subunits of APC/C, the multi-subunit RING finger E3 ubiquitin ligase that regulates exit from mitosis through the degradation of various mitotic regulators such as cyclin B and Securin (Carroll and Morgan, 2002; Kominami et al., 1998; Passmore et al., 2003). Here, we showed that Sox21 and Anapc10 negatively regulate Smad3 activity. Upon being relieved from this regulation, the increased Smad3 activity led to a potentiated TGF response toward EMT in the dental epithelium. Anapc10 has been reported to form a substrate recognition module together with Fzr1 or Cdc20 to recognize the D-box motif present in substrates for ubiquitination and subsequent degradation (Kominami et al., 1998). Our results suggest that Anapc10 and Fzr1 positively regulate E-cadherin and negatively regulate Smad3, N-cadherin, and Skp2 at the protein level. Furthermore, Anapc10 directly binds to Sox2 with ubiquitin-conjugating enzyme Ube2c and regulates Sox2 stability (Wang et al., 2016a). These observations suggest that Anapc10, together with Fzr1, prevents EMT in the dental epithelium and may regulate Sox2 expression. Our data further indicate that although Smad3 contains a consensus D-box motif sequence, it is unlikely that APC/C^{Fzr1} directly ubiquitinates Smad3 for degradation (Figures 5C, 5D, 6, S7C, and S7D). Instead, the previous observation (Nourry et al., 2004) that Anapc10 directly binds to Smad3 to trigger the ubiquitination of NEDD9/HEF1, a critical positive regulator of EMT, suggests that the EMT elicited by Sox21 KO in the dental epithelium may stem from a stabilization of NEDD9/HEF1 partly through decreased Anapc10 expression and impaired APC/C activity (Beck et al., 2014; Morimoto et al., 2014).

In addition, we present evidence that *Anapc10* knockdown led to Skp2 stabilization and the concomitant destabilization of E-cadherin in SF2 cells (Figure 6). As previous studies have demonstrated that APC/ C^{Fzr1} degrades Skp2 (Bashir et al., 2004; Wei et al., 2004) and that Skp2 directs E-cadherin degradation (Inuzuka et al., 2012), our data support the hypothesis that the Anapc10/Fzr1/Skp2/E-cadherin signaling pathway contributes to the EMT observed in *Sox21* KO dental epithelium. During tooth and hair follicle development, ectodermal cells form a bud structure by changing their polarity and cell-cell contacts via





E-cadherin. The transcription factor Lef1 downregulates E-cadherin to induce epithelial bud invagination. Moreover, forced elevation of E-cadherin expression blocks invagination and hair follicle production (Jamora et al., 2003). Overexpression of Lef1 under the control of the K14 promoter induced hair formation in the incisor region similar to our observation in Sox21-deficient mice (Zhou et al., 1995). These results suggest that downregulation of E-cadherin may be critical for tooth hair formation.

Taken together, our data suggest the existence of ameloblast plasticity that is maintained under the control of Sox21. Sox21 inhibited dental epithelial EMT and downregulation of E-cadherin via Anapc10. This information is useful for applications in the regeneration of organs including teeth and hair.

Limitations of the Study

This study reveals a role for Sox21 in ameloblasts. Inhibition of Sox21 switched the cell fate of dental epithelium to hair cells. Furthermore, the dental epithelium undergoes EMT and changes into mesenchymal cells. However, this may be limited to the incisors in rodents that continue to erupt. On the other hand, it may contribute to the elucidation of the differentiation of other ectodermal organs and may lead to the progress of the function of Sox21 expressed in the neural crest because tooth cells are derived from the neural crest.

Resource Availability

Lead Contact

Further information and requests for resources and reagents should be directed to and will be fulfilled by the Lead Contact, Kan Saito (kanta@dent.tohoku.ac.jp).

Materials Availability

This study did not generate new unique reagents.

Data and Code Availability

The data that support the findings of this study are available from the Lead Contact on reasonable request. Microarray data have been deposited in NCBI's Gene Expression Omnibus (GEO) and are accessible through GEO series accession number GSE99359 (https://www.ncbi.nlm.nih.gov/geo/query/acc.cgi? acc=GSE99359), GSE99360 (https://www.ncbi.nlm.nih.gov/geo/query/acc.cgi?acc=GSE99360).

METHODS

All methods can be found in the accompanying Transparent Methods supplemental file.

SUPPLEMENTAL INFORMATION

Supplemental Information can be found online at https://doi.org/10.1016/j.isci.2020.101329.

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AUTHOR CONTRIBUTIONS

K.S. and S.F. conceived and designed the whole experiments. F.M., H.O., and I.T. designed the experiments and interpret data. S.Y, K.Y, M. I., A.Y., E.F., S.Y., and T.N. collected samples and analyzed data. H.I. performed ubiquitination assay. K.S., F.M., and S.F. wrote the manuscript.

DECLARATION OF INTERESTS

The authors declare no competing interests.

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Supplemental Information

Sox21 Regulates Anapc10

Expression and Determines

the Fate of Ectodermal Organ

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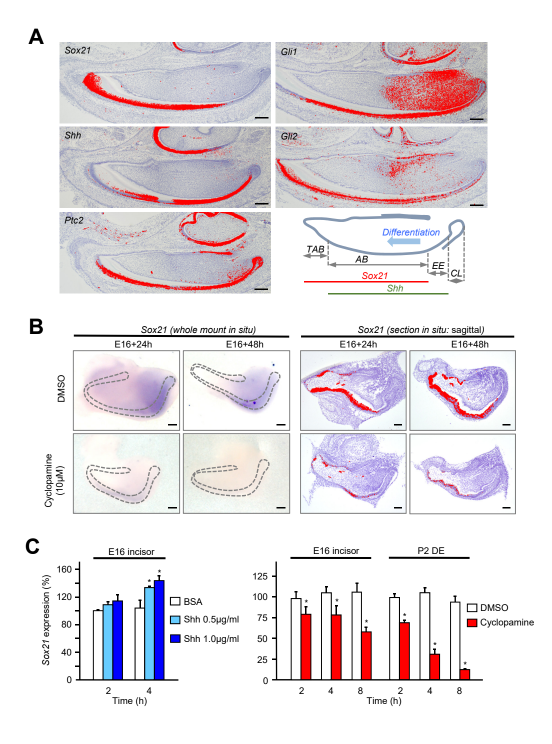


Figure S1. Expression and effect of Shh-related genes in the tooth. Related to Figure 1.

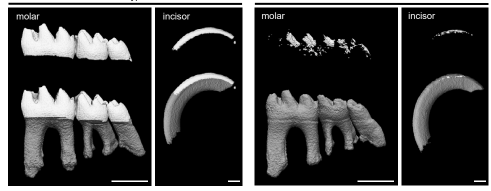
(A) Radioactive *in situ* hybridization on sagittal sections of the lower incisor from 6-week-old mice. ³⁵Slabeled RNA probes were used to detect the expression of *Sox21*, *Shh*, *Ptc2*, *Gli1*, and *Gli2*. Ptc2 of the receptor and Gli of the intracellular signal were used for analysis of Shh signaling. Scale bar, 200 μm. CL, cervical loop. EE, enamel epithelium. AB, ameloblasts. TAB, terminal ameloblasts.

(B) Incisors were dissected from E16-stage mice under a stereomicroscope. Tissues were cultured using a Trowell organ culture system for 24 or 48 h with the Shh inhibitor cyclopamine (10 μ M). Whole-mount *in situ* hybridization using digoxigenin-labeled probes was performed. Radioactive *in situ* hybridization on sagittal sections of the lower incisor was carried out in accordance with standard protocols. Scale bar, 200 μ m.

(C) E16 tooth germs or P2 dental epithelium (DE) were cultured with recombinant Shh or cyclopamine. Sox21 expression was quantified by qPCR and was normalized to that of Gapdh expression. The expression in the control sample was set to 100 and expression in the experimental samples was determined as the fold change relative to the control using the $\Delta\Delta$ CT method. Error bars represent mean ± SEM of five technical replicates. **p* < 0.05, Student's *t*-test. Α

wild-type

Sox21 KO



В

		Vol	ume (mm³)	
		wild-type	Sox21 KO	Fold change (<i>Sox21</i> KO / wild-type)
incisor	Enamel	0.1617349	0.0141242	0.0873293
	Dentin	1.8942210	2.1707440	1.1459824
molar	Enamel	0.3113039	0.0187767	0.0603163
	Dentin	1.2306690	1.2484150	1.0144198

Figure S2. Micro-computed tomography (micro-CT). Related to Figure 2.

Molars and incisors of 6-week-old wild-type and *Sox21* KO mice were analyzed by micro-CT. Mineralization volumes (mm³) of enamel and dentin were calculated. The tooth enamel and dentinal quantity were determined by the density of mineralization.

(A) Calcium scores of incisors and molars scanned by micro-CT. The lower figure is the whole tooth and the upper figure is the enamel. Scale bar, 1mm.

(B) Tooth enamel and the dentin-cementum were divided on the basis of the difference in calcification density, and the volumes were measured. Fold changes were calculated from the differences in density between wild-type (WT) and *Sox21* KO (KO) mice samples.

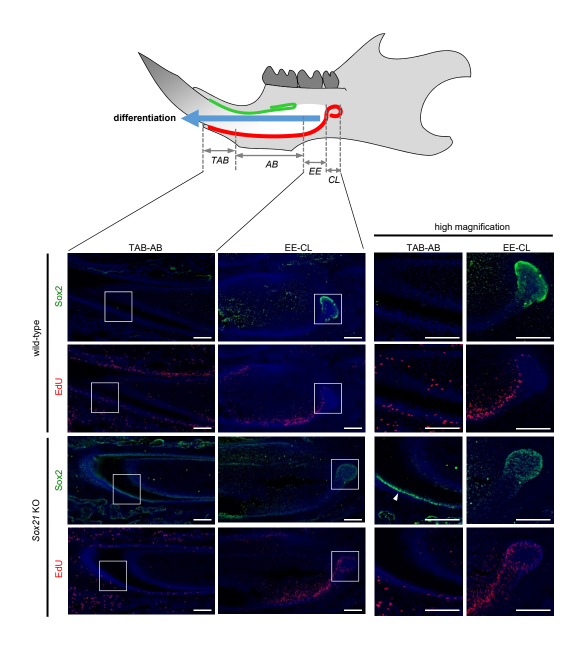
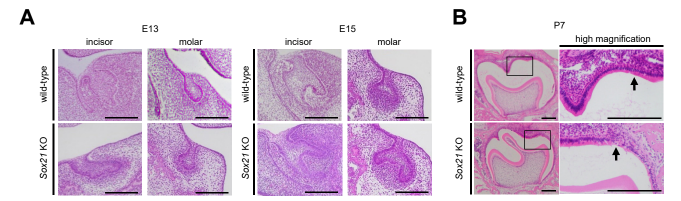
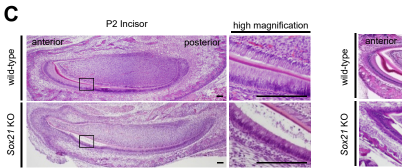


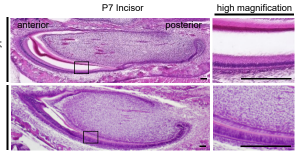
Figure S3. Sox2 expression and cell proliferation assay in *Sox21* KO mouse incisors. Related to Figure 3.

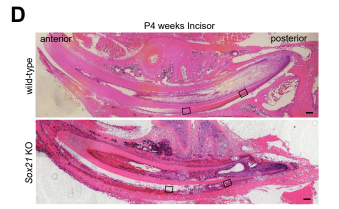
Schematic of a sagittal section through the mouse incisor. The cervical loop (CL) exists in the posterior region of the incisor. The enamel epithelium (EE) differentiates sequentially into ameloblasts (AB) and terminal ameloblasts (TAB), while cells migrate anteriorly from the labial CL. The red signal is EdU; green signal, Sox2 immunostaining. The right panel shows these results at high magnification. Scale bar, 200 μ m.

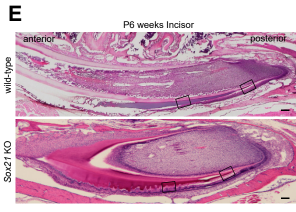
Figure S4

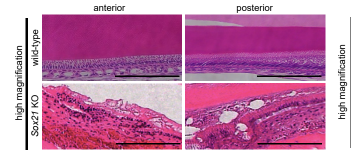












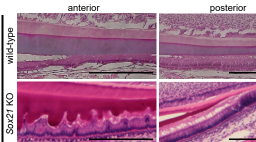


Figure S4. Histological analysis of the wild-type and Sox21 KO mice. Related to Figure 3.

(A) Incisors and molars in the embryonic stage of wild-type and *Sox21* KO mice were stained with hematoxylin and eosin (H-E). The left panel shows a sagittal section of the incisor; right panel, coronal section of the molar.

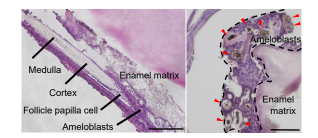
(B) The molar of 7-day-old wild-type and *Sox21* KO mice were stained with H-E. The right panel show high magnification of ameloblasts in a molar.

(C) The incisors of 2 and 7 days post-natal wild-type and *Sox21* KO mice were sectioned in a sagittal plane. The right panel shows these results at high magnification.

(D, E) The incisors of 4- and 6-week-old mice were sectioned in a sagittal plane. The lower panel shows these results at high magnification. Scale bar, 200 μ m.









Sox21 KO





В

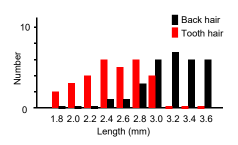


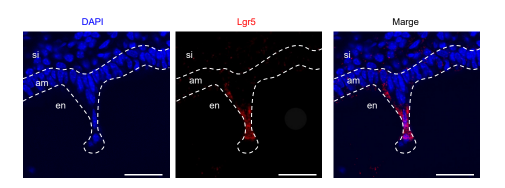
Figure S5. Anatomical observations of tooth hairs in *Sox21* KO mice. Related to Figure 3.

(A) For observation of tooth hairs, 5- and 6-week-old *Sox21* KO mouse lower incisor gingiva were assessed with a stereomicroscope. Arrow indicates hair in the gingiva. Scale bar, 1 mm.

(B) Histological analysis of the 6-week-old *Sox21* KO mouse lower incisor. Left panel shows sagittal section, right panel shows frontal section. Red arrowheads indicate the hair root sheath structure. Scale bar, 100 μ m.

(C, D) The tooth and back hairs of 6-week-old *Sox21* KO mice were examined by stereomicroscopy. Stereomicroscopy was used to investigate hair morphology (C). Scale bar, 1 mm. Thirty hairs on each tooth and back were measured (D).

Α



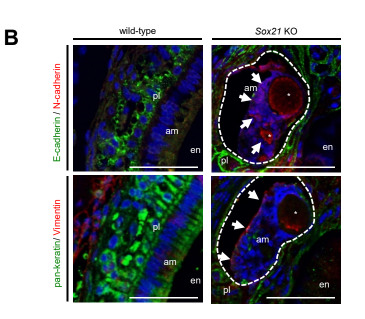
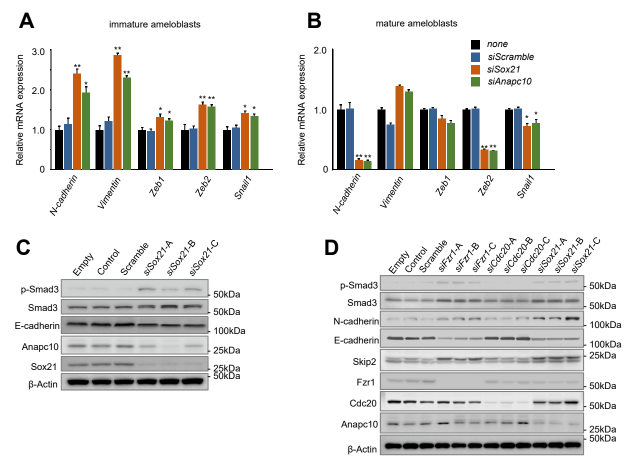


Figure S6. Immunohistochemistry of hair root sheath structures derived from odontogenic epithelium. Related to Figure 3.

(A) The ameloblasts of 6-week-old *Sox21* KO mouse incisors were immunostained using anti-Lgr5 antibody with DAPI. si, stratum intermedium; am, ameloblasts; en, enamel. Scale bar, 100 µm.

(B) Double immunostaining of incisal frontal section from a 6-week-old mouse, counterstained with DAPI. Upper panels show sections stained using anti-E-cadherin and anti-N-cadherin antibodies; lower panels, anti-pan-keratin and anti-Vimentin. Scale bar, 100 µm. am, ameloblast; en, enamel. pl, papillary layer; arrow, mesenchyme marker positive cells; *root sheath.



Ε

		siScra	amble	
	E-cadherin	β-Actin	DAPI	Phase
none				
TGF-β1				

		siCc	lc20	
	E-cadherin	β-Actin	DAPI	Phase
none				
TGF-β1	and a			

Figure S7. Induction of EMT by inhibition of Sox21 related genes. Related to Figure 4.

(A, B) Sox21 or Anapc10 expression in SF2 cells was repressed using siRNA. Scramble siRNA was used as a negative control. After 48 h, the expression of *N*-cadherin, Vimentin, Zeb1, Zeb2, and Snail1 was evaluated by qPCR. SF2 cells were used as rat immature ameloblasts (A). SF2 cells were cultured with NT-4 for differentiation to mature ameloblasts (B). Error bars represent mean \pm SEM of five technical replicates. Student's *t*-test (**p* < 0.05, ***p* < 0.005).

(C, D) SF2 cells were cultured with the indicated siRNAs for 48 h, harvested in cell lysate buffer with a proteinase inhibitor mixture, and subjected to western blotting. SF2 cells transfected with scramble and three different types (-A, -B, -C) of *Sox21* siRNA (C). SF2 cells were cultured with *Fzr1*, *Cdc20*, and *Sox21* siRNAs (three types for each) (D).

(E) SF2 cells were transfected with scramble or *Cdc20* siRNA probes. Transfected cells were cultured with 1 ng/mL TGF-β1 for 48 h. Fixed cells were incubated with anti-E-cadherin (green) and anti-β-Actin (red) primary antibodies. Expression was detected using Alexa 488- or Alexa 594-conjugated secondary antibodies. Cells were counterstained with DAPI (aqua). Fluorescence microscopy was used for image analysis. Images were prepared using a BZ analyzer.

Table S1

				P1 Sox21KO P1 WT tooth germ and WT tooth germ and skin			1	
			wild-type	Sox21 KO	Sox21KO /WT	Tooth	Skin	Skin / Tooth
	Gene	D	normalized	normalized	foldchange	normalized	normalized	fold change
	Wht16	NM_053116	0.545214	0.110121	0.2019776	0.07893681	1.01384586	12843766
	Stc24a4	NM_172152	1.485913	0.307624	0.2070268	0.39510026	0.35461334	0.8975275
ç	Nfasc	NM_182716	35188	0.899331	0.2555788	0.77622677	0.1611484	0.2076048
atic	Amin	NM_027793	13.05109	3.526093	0.2701761	1.9298519	0.0180962	0.009377
Down regulation	Gdf5	NM_008109	7.573416	2.282076	0.3013271	1.99269568	0.61521532	0.3087352
L L	Gal	NM_010253	4.820243	1.654346	0.343208	6.11920868	2.15714656	0.3525205
Po	Lama3	NM_010680	2229153	8233238	0.3693437	18.1096405	5.62769133	0.3107567
	Wni8b	NM_011720	0.72575	0.345303	0.4757878	025213238	0.07998837	0.3172475
	Lamb3	NM_008484	70.09956	33.37616	0.4761251	46.3924779	16.993905	0.3663073
	Klk4	NM_019928	46.50384	22.53712	0.4846291	10.476001	0.17508592	0.016713
	Kn6a	NM_008476	4.328279	10.50092	2426119	5.04476617	3.98179156	0.7892916
	Sox2	NM_008476	0.084384	0.204748	2.4263943	0.06794838	0.10708674	1.5760013
	Serpinb2	NM_011111	0.09823	0.241208	24555493	0.06468927	2.66754757	41.236323
	Eppk1	BC026387	2439613	5.99995	24593853	5.0134961	43.1576469	8.6082937
	<i>lii202</i> b	NM_011940	020634	0.509715	24702613	2.38397697	0.61965206	0.2599237
	Aqp3	NM_016689	4.686664	11.58602	24721252	224232905	432737428	19.298569
	Cabr	NM_007588	0.477696	1.230294	25754752	0.01744918	0.01597048	0.9152569
	Gjb3	NM_001160012	0.463193	1.207677	2.6072894	0.93076087	50.1520337	53.882834
	Gjb2	NM_008125	0.434406	1.133406	2.6090954	0.68838086	25.6058581	37.197225
	Dsg3	NM_030596	0.361908	0.948636	2621204	0.37224052	27124189	7.2867374
tion	Knt6b	NM_010669	4.638635	1217714	2625157	238215898	7.8478616	3.2944323
Up regulation	Scel	NM_022886	0.341166	0.926424	27154678	0.84052962	10.2047208	12.140822
o reç	Kit1	NM_008473	0.73345	2.006899	27362462	0.56873695	563.543826	990.86902
5	Mais2	NM_001159569	0.540254	1.563286	28936115	0.32500126	0.9318031	28670754
	K123	NM_033373	1269915	3.877613	3.053442	1.16792106	59.0901477	50.5943
	K i k13	NM_001039042	0.150493	0.469854	3.1220977	0.13142733	4.63268086	35.248992
	Kit15	NM_008469	1211519	39.41645	32534733	9.01582555	215.751428	23.930302
	Calm4	NM_020036	1.18064	4.17527	3.5364453	1.32356009	677.933992	51220492
	Lgr5	NM_010195	0.147347	0.52749	3579916	0.24952161	3.94997083	15.830175
	Kit10	NM_010660	0.409956	1.470797	3.5876899	2.55683595	871.239769	340.74919
	Kit13	NM_010662	3.106192	11.88748	3.8270265	11.6189001	602434714	5.1849548
	Spink5	NM_001081180	0.677767	2765247	4.0799382	0.67279899	29.5925682	43.98426
	Ktt32	NIM_001159374	0.113465	0.597378	52648459	0.1467483	0.75555202	5.148625

Table S1. Microarray analysis of ameloblasts in the Sox21 KO mice Related to Figure 3.

Comparing wild-type (WT) and *Sox21* KO (KO), the molecules with decreased expression are shown in blue, and the molecules with increased expression are shown in red. Comparing skin and tooth of wild-type mice, the Down-regulation molecules are shown in blue, and the Up-regulation molecules are shown in red.

Table S2

Primers for qPCR

Gene	Forward	Reverse
Gapdh	TCCTGCACCACCAACTGCTTA	AGTGATGGCATGGACTGTGGT
Sox21	GGGGCCCGGTTTGTATGT	CAACTGCTGGCGCTAACAAC
Slc24a4	TGATCACCAACAAGTTTGGGC	GCTGTGGGTCTTCAGGGTTC
GDF5	AAAGGGCAAGATGACCGAGG	TAAGATCCGCAGTTCAGCCC
Amtn	CCTTATCCACCCCTTGTTCCC	TCTGTGACASCAGGAGTTGTG
Lamb3	GGAGTCAGAGCTGCTTCGAG	GTAACTGCGGATCTGCTCCA
Klk4	AGCAGCCGGATCATACAAGG	TCTTCTGAGAACAGTGCCGC
Krt24	GTAAGGGGTGGATTCGGAGC	GAGAAGGCCCCCATCATAGC
Krt6	GGAAATTGCCACCTACAGGA	GGTGGACTGCACCACAGAG
Sox2	AATCACAACAATCGCGGCGG	CTGGCGGAGAATAGTTGGGG
Klk13	CAGTGCGCCAACATTGAACT	GCCGTAGAGTTTGCCATTGC
Krt23	AGGATGGCAGTGGATGACTTC	CCTCCTGTTCCAGGTCTGT
Krt1	GCAAACTCAAATCAGCGAAACC	TTGCTCCTCTGGCAAATGCT
Lgr5	GCGTCTTCACCTCCTACCTG	GCTCCCTTGGGAATGTGTGT
Krt32	GTGAGGCAGCTGGAGAAAGA	TCAGGACACATGGTCAGCAC
Krt15	GGCCAGGTCAATGTGGAAATG	TAGACGCCACCTCCTTGTTC
Krt13	CTGGCATTGATCTGACCCGT	CCCTCCGATTCTTCTCTGCC
Krt10	GTGCAGCTCTCCCAGATTCA	TGTTGGTACTCGGCGTTCTG
Kradap	TCCTGAACTGGCACGTCAT	CGGCACTTCTCAGTCCTTTC
Anapc10	TCAGGCTGTYTGGTCACTCT	AGCCATCRGATTGCCAGTAA
N-cadherin	ACAAAGGCAGAAGAGAGACTGG	ATGAAGATGCCCGTTGGAGG
Vimentin	AGGCCGAGGAATGGTACAAG	GACTGCACCTGTCTCCGGTA
Zeb1	CAGGAGGAGCCCCAAGTAGAA	GTTGGCACTTGGTGGGACTAC
Zeb2	GTTGTGATCCTCCTCTCAGGC	TAATGACAGGTCCAACGGCT
Snail1	TCACCTTCCAGCAGCCCTAC	TTGCCACTGTCCTCATCGGA

Table S2. Primer sequences Related to Figure 3.

These primers were used with SYBR Select Master Mix and Step One Plus Real-Time PCR system.

Transparent Methods

KEY RESOURCES TABLE

REAGENT or RESOURCE	SOURCE	IDENTIFIER
Antibodies		
Anti-Sox2	Abcam	Cat# Ab137385
Anti-Ameloblastin	Santa Cruz	Cat# sc-50534, 33100
Anti-Pan-keratin	ThermoFischer	Cat# 18-0059
Anti-Vimentin	GeneTex	Cat# GTX85471
Anti-E-cadherin	R&D systems	Cat# AF748
Anti-N-cadherin	GeneTex	Cat# GTX112734
Anti-β-Actin	Abcam	Cat# mAbcam8226
Anti-Alexa488	ThermoFischer	Cat# A11055/A21206
Anti-Alexa594	ThermoFischer	Cat# A11058/A21207
Anti-Smad3	Cell Signaling Technology	Cat# 9513
Anti-phospho Smad3	Cell Signaling Technology	Cat# 9520
Anti-Smad6	Sigma-Aldrich	Cat# WH0004091M7
Anti-Smad7	Sigma-Aldrich	Cat# SAB2108469
Anti-Sox21	LifeSpan BioSciences	Cat# LS-C107486
Anti-Fzr1	Santa Cruz	Cat# sc56312
Anti-CDC20	Santa Cruz	Cat# sc13162
Anti-Anapc10	Atlas Antibodies	Cat# HPA044547
Anti-Skp2	Santa Cruz	Cat# F1714
Anti-DDK	Origene	Cat# TA50011
Chemicals, Peptides, and Recombinant Prote	ins	
Cyclopamine	Merck Millipore	Cat# 239803
Sonic hedge hog	R&D Systems	Cat# 464-SH
TGF-β1	R&D systems	Cat# 240-B
NT-4	Alomone Labs	Cat# N-270
cycloheximide	Sigma-Aldrich	Cat# C1988
Superscript Vilo MasterMix	ThermoFischer	Cat# 11755050
SYBR® Select Master Mix	Applied Biosystems	Cat# 4309155
ViaFect™ Transfection Reagent	Promega	Cat# E4981
Critical Commercial Assays		
miRNeasy mini kit	QIAGEN	Cat# 217004
ChIP-IT Express kit	Active Motif	Cat# 53008
ECL Prime kit	GE Healthcare	Cat# RPN2232
Deposited Data		
RNA-seq data of Sox21KO mouse	This paper	GSE99359
RNA-seq data of skin and tooth	This paper	GSE99360
Experimental Models: Cell Lines		
•		
SF2 cells	Arakaki et al., 2012	N/A

Experimental Models: Organisms/Strains			
C57BL/6J mice The Jackson Laboratory JAX: 0			
K14-Follistatin mice	Wankell et al., 2001	N/A	
follistatin knockout mice	Matzuk et al., 1995	N/A	
Sox21 knockout mice	Kiso et al., 2009	N/A	
Recombinant DNA			
pCRII-TOPO	ThermoFischer	Cat# K460001	
pEF6/V5-His TOPO	ThermoFischer	Cat# K961020	
pCMV6-Entry	Origene	Cat# PS100007	
Sequence-Based Reagents			
Primers for Sox21 probe; Forward GATGTATAGGTGTCAGGCA	This paper	N/A	
Primers for Sox21 probe; Reverse GGTCATTCACTGGTCAAG	This paper	N/A	
Primers for qRT-PCR, see Table S2	This paper	N/A	
Software and Algorithms			
integrative genomics viewer	http://software.broadinstit ute.org/software/igv/	N/A	
ImageQuant	GE Health Life Science	N/A	
ImageJ	https://imagej.nih.gov/ij/	N/A	
Photoshop	adobe	Ver.11.0	

EXPERIMENTAL MODEL AND SUBJECT DETAILS

Animals

C57BL/6J mice were used as wild-type (WT) mice. *Keratin 14* promoter-driven ectopic follistatin gene expression (*K14-Follistatin*), follistatin knockout, and *Sox21* knockout (*Sox21* KO) mice were gifted. All animal experiments were approved by the Animal Ethics Committee of Tohoku University. *Sox21* KO mice were provided with a soft diet. These mice were euthanized by cervical dislocation under isoflurane anesthesia for all experiments. All experiments were performed in accordance with the Law Concerning the Conservation and Sustainable Use of Biological Diversity through Regulations on the Use of Living Modified Organisms. All animal research and genetic recombination experiments were approved by the Tohoku University Center for Gene Research (2013DnLMO-008, 2013DnA-051).

METHOD DETAILS

SEM Analyses

The heads of 6-week-old mice were stripped of the skin and treated at 48°C in 0.3% trypsin buffer in one day. Soft tissue was removed from the skull and the molars and incisors were extracted. Teeth were embedded in epoxy resin (Oken Epok; Okenshoji Co., Ltd.). Resin was cut using a diamond disk, cleaned ultrasonically after being polished, and examined using a variable pressure scanning electron microscope

(Miniscope TM3000; Hitachi). Elemental mapping at the microstructural level was investigated by SEM with an energy dispersive X-ray spectrometry system (Quantax70; Bruker).

Micro-CT Scanner Imaging

The skulls of 6-week-old wild-type and *Sox21* KO mice were analyzed by micro-CT at the Kureha Special Laboratory (Iwaki, Japan). The mandibles were imaged and the mineralization volumes (mm³) of enamel and dentin were calculated from the tomographically sliced sections. The tooth enamel and the dentinal quantities were determined by the density of mineralization.

Preparation of Tissue Sections and H-E Staining

Mice from E15 to post-natal 6 weeks were dissected in Dulbecco's PBS. The mandibles from the embryonic stages were fixed by 4% paraformaldehyde for 1 week. Half mandibles of P2 to post-natal 6-week-old mice were fixed and decalcified by using 2.5% paraformaldehyde and 12.5% EDTA in PBS for 2-6 weeks. The tissues were then dehydrated and embedded in paraffin. Paraffin-embedded sections (7-µm-thick) were prepared using a Leica microtome. For staining, sections were re-hydrated and stained with H-E (J. T. Baker).

In Situ Hybridization and Immunostaining

A Sox21 probe fragment comprising mouse *Sox21* mRNA [NM_177753] (nucleotides 2331–2825; 495 bp) was amplified using 5'-GATGTATAGGTGTCAGGCA-3' and 5'-GGTCATTCACTGGTCAAG-3' primers and cloned into pCRII-TOPO (Invitrogen). The fragment sequence was verified by sequencing. Radioactive *in situ* hybridization on 7-µm paraffin frontal and sagittal sections was carried out in accordance with standard protocols (Juuri et al., 2012). ³⁵S (Amersham)-labeled RNA probes were used for detection. Antigen was activated using Lab solution (Polysciences, Inc.) at 22°C for 15 min. Sections were incubated in 5% bovine serum albumin/PBS for 1 h prior to incubation with the primary antibody. Primary antibodies including anti-Sox2 antibody (1:100, Abcam), anti-Ameloblastin (1:50, Santa Cruz), anti-pan-keratin (1:100, Thermo Fischer Scientific), anti-Vimentin (1:100, GeneTex), anti-E-cadherin (1:20, R&D Systems), and anti-N-cadherin (1:100, GeneTex) were detected using Alexa488- or Alexa594-conjugated secondary antibodies (Thermo Fisher Scientific). Nuclei were stained with DAPI (Vector Laboratories). A fluorescence microscope (BZ-8000; Keyence) and confocal microscope (Fluoview Fv10i LIV; Olympus) were used for imaging analysis. Images were prepared using Adobe Photoshop (Adobe Systems, Inc.).

In vivo EdU staining

Edu (50 mg/kg) in PBS was intraperitoneally injected for 2 days postnatal mice. Mice were dissected after 2 h, the lower mandibles were frozen with O.C.T. Compound (Sakura Finetek Japan). Sections (20-µm-thick) were prepared using a Leica cryostat. EdU was detected in accordance with the protocol of EdU proliferation kit (Abcam). Counterstaining of Sox2 and DAPI were performed by Anti-Sox2 antibody and mounting medium with DAPI.

Cell Culture and siRNA Treatment

SF2 (rat immature dental epithelial cell line) was cultured in DMEM/F12 medium (Life Technologies) containing 10% fetal bovine serum (Life Technologies) and an antibiotic cocktail (Gibco, NY, USA) at

37°C and 5% CO₂. Mature ameloblasts differentiate from dental epithelial cells with Neurotrophin (NT)-4 (Yoshizaki et al., 2008). SF2 cells were cultured with 100 ng/mL recombinant human NT-4 (Alomone Labs) for differentiation to mature ameloblasts. For inhibition of *Sox21*, *Anapc10*, *Fzr1*, and *CDC20* expression, siRNAs were transfected into cells using Stealth RNAi[™] (Thermo Fisher Scientific) with Lipofectamine[™] RNAiMAX (Thermo Fisher Scientific) in accordance with the manufacturer's protocol. Cells were harvested after 48 h.

Expression Vector and Transfection

The *Sox21* open reading flame was amplified from P2 mouse incisors (5'-ATGTCCAAGCCTGTGGACCACG-3', 5'-TAGCGCGGCAGCGTAGGC-3'). The PCR fragment was cloned into the pEF6/V5-His TOPO vector (Thermo Fisher Scientific). Full length *Sox21* sequence was verified by sequencing. SF2 cells were trypsinized and replated in a 6-well plate at a density of 1 × 10⁵ cells/well. The cells were transfected with plasmid DNA using the ViaFectTM transfection reagent (Promega), which was mixed in the culture plate at a 3:1 (ViaFectTM/DNA) ratio. The cells were collected for the experiments after 48 h.

RT-qPCR

Total RNA was extracted using the Quiagen RNase Kit in accordance with the manufacturer's instructions. Extracted RNA was converted into cDNA using Superscript Vilo MasterMix (Thermo Fisher Scientific). PCR was performed with the SYBR Select Master Mix (Applied Biosystems) and each specific primer (Table S2) pair using a Step One Plus Real-Time PCR system (Applied Biosystems).

Microarray Analysis

Skin and tooth germs were isolated from P1 wild-type (WT) and Sox21 KO (KO) mice. Labeled RNA samples were hybridized onto SurePrint G3 Mouse GE 8×60K microarrays (Agilent Technologies). Microarray data were analyzed using GeneSpring (Agilent Technologies). Total detected entities were filtered by signal intensity value to remove very low signal entities. Normalization was performed using a percentile shift in the analysis of signaling pathways. Therese data have been deposited in NCBI's Gene Expression Omnibus (GEO) and are accessible through GEO series accession number GSE99359, GSE99360.

ChIP-Seq

SF2 cells were transfected with the *Sox21* pCMV6-Entry vector (Origene) using ViaFect in accordance with the manufacturer's instructions. The medium was changed 24 h after transfection. After 48 h, 5 ×10⁶ cells were fixed by formaldehyde and processed using the ChIP-IT Express kit (Active Motif). Cells were lysed and then sonicated to shear the chromatin. The fragmented genomic DNA (5 μ g) was immunoprecipitated using the anti-DDK monoclonal antibody (Origene) overnight at 4°C. The ChIP reactions were washed and chromatin was eluted in accordance with the manufacturer's instructions. The 1 ×10⁷ DNA sequencing, computational, and statistical analyses of ChIP-Seq data were performed using Active Motif. The ChIP-Seq results were visualized using integrative Genomics Viewer (IGV) software (https://www.broadinstitute.org/igv/).

Western Blotting Analysis

SF2 cells transfected with *Sox21*, *Anapc10*, *Fzr1*, *Cdc20*, and Scramble siRNA (Sigma-Aldrich) were plated in 6-well plates at a concentration of 1 × 10⁵ cells per well and incubated for 48 h. The scramble sequence siRNA was using as the negative control. For Smad cell signaling analysis, the cells were then cultured without serum-containing medium for 2 h, followed by treatment with 1 ng/mL TGF-β1 for 0, 5, 15, 30, or 60 min at 37°C. Thereafter, the cells were washed twice with ice-cold 1 mM sodium orthovanadate (Sigma-Aldrich) in PBS, lysed with Nonidet P-40 buffer supplemented with a proteinase inhibitor mixture (Sigma-Aldrich), and centrifuged. The supernatants were then transferred to a fresh tube. For protein half-life analysis, siRNA treated cells were incubated with 300 µg/mL CHX for 0, 1, 2, and 4 h. The cells were then harvested by cell lysate buffer with a proteinase inhibitor mixture and moved to a fresh tube. The cell lysates were separated by 4-12% gradient SDS-PAGE and analyzed by western blotting. The blotted PVDF membrane was incubated with Smad3 (Cell Signaling Technology), Smad6 (Sigma-Aldrich), Smad7 (Sigma-Aldrich), or phosphor-Smad3 (Cell Signaling Technology) primary antibodies. The signals were detected using an ECL Prime kit (GE Healthcare) after treatment with a rabbit or mouse horseradish peroxidase-conjugated secondary antibody. Images were visualized using the ImageQuant LAS 4000 Mini image analysis system (GE Healthcare).

QUANTIFICATION AND STATISTICAL ANALYSIS

Relative Gene Expression Analysis by qPCR

The expression of each gene was normalized to that of *Gapdh* expression. The expression in the control sample was set to 1.0 and expression levels in the experimental samples were determined as the fold change relative to the control using the $\Delta\Delta$ CT method (n = 5). Statistical analysis of gene expression was performed using the Student's t test, with *p* < 0.05 considered significant.

Analysis of Cell Localization of E-cadherin and β-Actin

A line of 1500 pixels was selected from E-cadherin and β -Actin immunofluorescence images. The fluorescence intensity was measured using ImageJ software and graphed.

Data and Software Availability

In situ hybridization and immunohistochemistry photographs were merged using Adobe Photoshop. Meta data was aligned using IGV software (http://software.broadinstitute.org/software/igv/). Signal intensity was analyzed using ImageJ software (https://imagej.nih.gov/ij/). Western blotting was visualized with ImageQuant (GE Healthcare).