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Local application of *Usag-1* siRNA can promote tooth regeneration in *Runx2*-deficient mice

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Runt-related transcription factor 2 (*Runx2*)-deficient mice can be used to model congenital tooth agenesis in humans. Conversely, uterine sensitization-associated gene-1 (*Usag-1*)-deficient mice exhibit supernumerary tooth formation. Arrested tooth formation can be restored by crossing both knockout-mouse strains; however, it remains unclear whether topical inhibition of *Usag-1* expression can enable the recovery of tooth formation in *Runx2*-deficient mice. Here, we tested whether inhibiting the topical expression of *Usag-1* can reverse arrested tooth formation after *Runx2* abrogation. The results showed that local application of *Usag-1* Stealth small interfering RNA (siRNA) promoted tooth development following *Runx2* siRNA-induced agenesis. Additionally, renal capsule transplantation of siRNA-loaded cationized, gelatin-treated mouse mandibles confirmed that cationized gelatin can serve as an effective drug-delivery system. We then performed renal capsule transplantation of wild-type and *Runx2*-knockout (KO) mouse mandibles, treated with *Usag-1* siRNA, revealing that hindered tooth formation was rescued by *Usag-1* knockdown. Furthermore, topically applied *Usag-1* siRNA partially rescued arrested tooth development in *Runx2*-KO mice, demonstrating its potential for regenerating teeth in *Runx2*-deficient mice. Our findings have implications for developing topical treatments for congenital tooth agenesis.

Tooth anomalies are common congenital anomalies in humans, who show a high incidence of missing teeth and affects one percent of people world-wide¹. Cases of more than six missing teeth can be caused by genetic factors⁶, and inherited tooth agenesis occurs in 10% of all individuals with congenital tooth agenesis⁶. Several different gene mutations have been identified in patients with congenitally inherited tooth agenesis⁶. In many cases, candidate genes were identified and determined, based on phenotypic changes observed in knockout (KO) mice. A recent study of runt-related transcription factor 2 (*Runx2*)−/− mice showed that they exhibited arrested tooth development, and this phenotype was compared with that of a patient harboring a unique Arg131Cys missense *Runx2* mutation who exhibited congenitally missing tooth without supernumerary teeth⁷. The findings suggested that *Runx2*−/− mice may be suitable as murine models of congenital tooth agenesis.

Uterine sensitization-associated gene-1 (*Usag-1*) is expressed in odontogenic epithelial cells and is a common antagonist of bone morphogenetic proteins (BMPs) and members of the Wnt-signaling pathway. *Usag-1*-deficient mice exhibit supernumerary tooth formation⁸, and we previously demonstrated that arrested tooth formation was restored by crossing *Runx2*−/− mice with *Usag-1*−/− mice (which represent a mouse model of supernumerary tooth formation)⁹. Additionally, previous results demonstrated that the number of teeth can be increased by topical administration of BMP-7⁹, indicating the possibility of forming an organic tooth by local manipulation.

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of a single target molecule. However, although a genetic link between Usag-1 and Runx2 has been clearly shown, it remains unclear whether topical inhibition of Usag-1 expression can recover tooth formation in Runx2−/− mice.

In recent years, oligonucleotide therapies for use in clinical applications have attracted increased attention10–14, due to a higher specificity for their target molecules (i.e., mRNAs) than conventional drugs. These therapies include antisense RNAs, small-interfering (si)RNA, and RNA aptamers. Specifically, siRNAs promote RNA interference (RNAi) and represent next-generation candidate drugs based on their direct and specific abilities to target disease-related genes15,16. We previously demonstrated that Stealth siRNAs, which are chemically modified siRNAs, were effective at gene targeting in mHAT9d cells, a dental epithelial stem cell line derived from the labial cervical loop epithelium of mouse incisors17. Therefore, in the present study, we used Usag-1 Stealth siRNA to inhibit local expression.

In this study, we tested the hypothesis that topical inhibition of Usag-1 expression would rescue arrested tooth formation. We performed rescue experiments using Stealth siRNAs against Usag-1 and Runx2, which were administered using cationized gelatin as a drug-delivery system (DDS).

Results
Efficient tooth development following local application of Usag-1 and Runx2 Stealth siRNA. We confirmed the expression of Usag-1 in mHAT9d cells derived from mouse odontogenic epithelial cells (Fig. 1A) and designed two types of Usag-1 Stealth siRNAs (#304 and #903) (Table 1). In our previous study, we determined the necessary conditions for transfecting mHAT9d cells with Stealth siRNAs17. To evaluate the knockdown efficiency of Usag-1 siRNA, we performed semiquantitative reverse transcription–polymerase chain reaction (sqRT-PCR) analysis in mHAT9d cells, finding that Usag-1 mRNA levels were reduced by ~50% with Usag-1 Stealth siRNAs #304 and #903 (Fig. 1A). To evaluate the effect of Usag-1 knockdown on tooth development, we investigated mandible explant cultures using serum-free medium containing Stealth siRNA. We confirmed the same efficiency of Usag-1 knockdown by both #304 and #903 in mandible explant cultures relative to that observed in mHAT9d cells (Fig. 1B). After a 10-day culture, we investigated tooth development with respect
to staging and the number of tooth germs following transfection with both Usag-1 siRNAs and Runx2 Stealth siRNA #1623, the silencing efficacy of which was confirmed previously. We found comparable staging and a high number of tooth germs following Usag-1 knockdown and comparable staging and a low number of tooth germs following Runx2 knockdown (Fig. 1C). These results demonstrated the biological effects of local application of Usag-1 and Runx2 Stealth siRNA on tooth development.

Cationized gelatin hydrogel as a DDS for Stealth siRNA during renal capsule transplantation of mouse mandibles. To confirm the efficacy of cationized gelatin as a DDS for local administration of siRNA, we performed renal capsule transplantation of mouse mandibular explants combined with cationized gelatin delivery of siRNA (Fig. 2A). We performed renal capsule transplantation of the mandibles into wild-type mice as controls. On day 19 post-transplantation, the tissue attached to the kidney was removed. Based on histochemical staining, we observed that the sections contained tooth structures, although we found no differences in the results with or without cationized gelatin. Based on histochemical staining, we observed that the sections contained tooth structures, although we found no differences in the results with or without cationized gelatin. (C) Histological sections at days 3, 6, and 10 post-transplantation. Sections were evaluated by immunostaining. Magnification, 200×. Cells were immunostained with an Alexa Fluor 488-conjugated antibody.

Table 1. Sequences of Usag-1 siRNAs.

| Gene name | Sense | Anti-sense | NCBI accession number |
|-----------|-------|------------|-----------------------|
| USAG-1 knock down Stealth siRNA | | | |
| #304 | UCAUGACUCUGAGACUUGGAUCGAAA | UUUCGAUCCAGUCUCAGUCAGCUCAUGA | NM_025312 |
| #903 | CAAGUGUCUAAGAUGAAUGAGUA | UCUCUAAUACAUUGAGACACUCUG | NM_025312 |

Figure 2. Renal capsule assays using lateral mandibles from wild-type mice. (A) Schematic illustration of the procedure used for renal capsule transplantation. M indicates the lateral mandibles of E10 mice, G indicates cationized gelatin containing Stealth siRNA, and A indicates agarose agar (used to maintain the graft space). (B) We evaluated H&E-stained sections, where the lateral mandible from wild-type mice was transplanted beneath the kidney capsule of nude mice (KSN/Sli). Renal capsule assays were performed with two groups: lateral mandibles transplanted with or without a cationized gelatin sheet containing PBS. Based on histochemical staining, we observed that the sections contained tooth structures, although we found no differences in the results with or without cationized gelatin. (C) Histological sections at days 3, 6, and 10 post-transplantation. Sections were evaluated by immunostaining. Magnification, 200×. Cells were immunostained with an Alexa Fluor 488-conjugated antibody.
that cationized gelatin demonstrated controlled release of siRNA, suggesting its potential efficacy as a DDS for local siRNA administration in renal capsules transplanted with mouse mandibles.

**Usag-1 knockdown rescued tooth formation attenuated by Runx2 knockdown.** To evaluate the biological effect of local application of Usag-1 and Runx2 Stealth siRNA with cationized gelatin hydrogels on tooth formation, we performed renal capsule transplantation of wild-type mouse mandibles. The number of tooth structures was determined by histological evaluation of serial H&E sections and 3D micro-CT analysis. The number of tooth structures was confirmed by 3D reconstruction of serial H&E sections in representative samples (Fig. 3). We observed the formation of more than three tooth structures following treatment with both Usag-1 Stealth siRNA #304 and #903, whereas no more than two tooth structures were observed in the negative-control group (Fig. 3). Additionally, the incidence of no tooth formation following Runx2 Stealth siRNA treatment (66.7%) was higher than that in the negative-control group (33.3%) (Fig. 3). These results suggested that Usag-1 Stealth siRNAs increased the number of tooth structures formed, whereas Runx2 Stealth siRNA hindered tooth formation. To investigate whether administration of Usag-1 Stealth siRNAs could rescue the phenotype of inhibited tooth formation caused by Runx2 knockdown, we co-administered Usag-1 and Runx2 siRNA, revealing that Usag-1 #304 recovered tooth formation in the presence of Runx2 Stealth siRNA (P=0.020), whereas Usag-1 #903 administration resulted in a similar incidence of no tooth formation as observed with Runx2 knockdown alone (P=0.656) (Fig. 3). These results demonstrated that hindered tooth formation induced by Runx2 knockdown was rescued by Usag-1 siRNA #304 treatment in renal capsules transplanted with wild-type mouse mandibles.

**Topical application of Usag-1 Stealth siRNA partially rescued arrested tooth development in Runx2−/− mice.** To investigate whether the local application of Usag-1 Stealth siRNA in cationized gelatin hydrogels can recover arrested tooth development in Runx2−/− mice, we performed renal capsule transplanta-
portion of mouse mandibles in Runx2−/− mice along with Usag-1 #304 and #903 siRNA. In the absence of Usag-1 siRNAs, we observed flattened explants and no growth of transplanted mandibles (Fig. 4A-a). However, growing explants were observed (42.3%) following topical application of Usag-1 #304 siRNA (Fig. 4A-b), but not Usag-1 #903 siRNA (Fig. 4B). Moreover, we observed no mineralized hard tissue, such as bone, dentin, or enamel, in the rescued mandibles via micro-CT analysis (Fig. 4A-c). Furthermore, histological investigation of growing mandibles treated with Usag-1 Stealth siRNA #304 showed no tooth structure; however, odontogenic epithelial-like cells regularly arranged as elongated rectangular cells with nuclear polarity were observed (Fig. 5A-a,b). sqRT-PCR analysis revealed that genes encoding the enamel-specific proteins, amelogenin and ameloblastin, were faintly expressed (Fig. 5B), and subsequent immunohistochemistry for amelogenin verified local expression in odontogenic epithelial-like cells (Fig. 5A-c,d). These results demonstrated that the topical application of Usag-1 Stealth siRNA #304 partially reversed the arrest of tooth development in Runx2−/− mice.

Discussion
We previously demonstrated that arrested tooth formation in Runx2−/− mice was restored by crossing them with Usag-1−/− mice; however, it remained unclear whether topical inhibition of Usag-1 expression would recover tooth formation in Runx2−/− mice. In the present study, two Usag-1 siRNAs (#903 and #304) were evaluated in mHAT9d cells and mandible explant cultures, followed by local application in a model of sub-renal capsule mandible implantation combined with a cationized gelatin sheet as a DDS (Fig. 2A). This model was suitable for evaluating the number of teeth formed after local siRNA administration, although the positions and shapes of the tooth germs developing inside the capsules were not typical. Our in vitro results showed that arrested tooth formation induced by Runx2 Stealth siRNA was rescued by Usag-1 Stealth siRNA #304 (P = 0.020) but not Usag-1 Stealth siRNA #903 (P = 0.656) (Fig. 3). The differences in the effects of these Usag-1 siRNAs may be explained by off-target effects resulting from differences in mRNA-expression levels between wild-type cells and those with Runx2 knockdown. Similar results were subsequently confirmed in vivo in Runx2−/− mice (Fig. 4). In a previous study, no effect of siRNA administration was observed in Toll-like receptor 7-KO mice, whereas in this study, Usag-1 Stealth siRNA #304 effectively recovered congenital tooth agenesis, both in vitro and in vivo.

RNAi relies on siRNAs that target mRNAs based on high sequence specificity, with their use holding great promise for developing therapeutics directed against targets otherwise not addressable using current methods.
However, problems exist with the local administration of siRNAs in vivo: it is difficult to transfer siRNAs into target cells, it is difficult to sustain the effects through increased biostability, and an appropriate DDS is needed for selective delivery. In the present study, we achieved RNAi using Stealth siRNAs, which are chemically modified to eliminate off-target effects. These siRNAs show higher efficiency knockdown of target mRNA expression, higher specificity, greater stability, and less cellular toxicity. Furthermore, we demonstrated that cationized gelatin can serve as a DDS to control the release of siRNAs for the local administration of siRNAs in mouse mandibles (Fig. 2). Additionally, cationized gelatin microspheres have been shown to enhance and prolong the anti-fibrotic effects of heat shock protein 47 (Hsp47) siRNA in a murine model of unilateral ureteral obstruction.

The United States Food and Drug Administration recently approved patisiran, an RNAi therapeutic agent, for the treatment of polyneuropathy in adult hereditary amyloid transthyretin amyloidosis. Molecularly targeted therapy involves the use of small molecules, monoclonal antibodies, or siRNAs to identify and target specific cells by interfering with target molecules. Additionally, this method induced de novo tooth formation via in situ repression or activation of a single candidate molecule in congenital tooth agenesis. In mouse and dog models of congenital tooth agenesis associated with ectodysplasin A (EDA) deficiency, administering an agonist antibody targeting EDA1 or recombinant EDA1 protein rescued the number, position, shape, and timing of the eruption of missing teeth. In the present study, we demonstrated that Usag-1 Stealth siRNA #304 effectively promoted tooth regeneration in Runx2−/− mice demonstrating arrested tooth development. This finding suggests the therapeutic potential of topical Usag-1 Stealth siRNA application via cationic gelatin for treating patients experiencing congenital tooth agenesis due to Runx2 deficiency. However, we found that topical Usag-1 Stealth siRNA application was insufficient for whole-tooth regeneration due to its limited potential in inhibiting Usag-1 mRNA expression (~ 50% in vitro; Fig. 1).

In conclusion, we confirmed the biological effectiveness of Usag-1 Stealth siRNA at rescuing arrested tooth development in Runx2−/− mice. Additionally, we demonstrated the effectiveness of cationized gelatin as a DDS for the local administration of Stealth siRNAs to promote tooth regeneration. Our findings suggest that the method developed in this study shows potential for future clinical applications aimed at treating patients with congenital tooth agenesis resulting from Runx2 mutation.

Figure 5. Immunohistochemical evaluation of serial sections and sqRT-PCR analysis from renal capsules transplanted with mandibles from E10 Runx2-KO mice and treated with Stealth siRNAs. (A-a, b) H&E staining showed no tooth structures; however, odontogenic epithelial-like cells with elongated, rectangular morphology were observed. Magnification: 200× and 400×. (A-c) Immunostaining for amelogenin was positive in odontogenic epithelial-like cells. Magnification: 400×. (A-d) Control. (B) sqRT-PCR analysis of the enamel-specific proteins, amelogenin and ameloblastin. sqRT-PCR at 19 days post-transplantation, using wild-type and Runx2-KO mouse mandibles with Usag-1 Stealth siRNA #304 or PBS.
Methods
Ethics statement. This study was approved by the Animal Research Committee of Kyoto University (approval number Med Kyō 19281) and the Recombinant DNA Experiment Safety Committee of Kyoto University (for performing recombinant DNA experiments). All experiments were performed in accordance with ARRIVE guidelines. All procedures were performed in accordance with relevant guidelines in the manuscript.

Mouse strains. Runx2−/− and wild-type mice were used in this study, both of which had a C57Bl6/J background. Embryos were obtained by timed mating, with day 0 (E0) considered to start at midnight on the day prior to detection of a vaginal plug. We used mice at day 10 (E10) for all experiments.

siRNA preparation. We designed sense and antisense Stealth RNAi siRNAs for Usag-1 and Runx2 (Table 1) and measured gene expression by qRT-PCR following transfection to evaluate target gene knockdown. mHAT9d cells were cultured on Primaria tissue culture plates (BD Biosciences, Franklin Lakes, NJ, USA)37. For siRNA transfection, we prepared Stealth siRNA–Lipofectamine RNAiMAX complexes using siRNA (6.0 pmol) (Stealth RNAi siRNA Negative Control, Med GC Dup, Stealth RNAi siRNA #304, or Stealth RNAi siRNA #903; Thermo Fisher Scientific, Waltham, MA, USA). For each well of a 24-well plate, the complexes were diluted in 100 μL Opti-MEM I reduced-serum medium and incubated at room temperature for 20 min. Then, the transfection cocktail was added to each cell suspension (500 μL) in growth medium and incubated at 37 °C in a CO2 incubator for 48 h. Total RNA was extracted from adherent cells using the TRizol reagent (Invitrogen, Carlsbad, CA, USA), chloroform, and phenol chloroform. RNA concentrations were measured using a NanoDrop2000 system (Thermo Fisher Scientific). Reverse transcription of total RNA was performed using a SuperScript IV first-strand synthesis system (Thermo Fisher Scientific). The synthesized complementary DNA (cDNA) samples were serially diluted (1/10, 1/30, and 1/90) in TE buffer and analyzed by PCR. PCR amplification was performed using the Ex Taq reagent (TaKaRa Bio, Shiga, Japan) or KOD FX (TOYOBO, Osaka, Japan) and specific oligonucleotide primers for the target sequences. Bands were quantitated with a bio-image analyzer (FAS-IV; Nippon Genetics, Tokyo, Japan).

Organ culture. E10 Runx2−/− and wild-type mouse mandibles were dissected under a stereomicroscope. Tooth explants were transfected with Stealth RNAi siRNA #304, #903, #1623, or a negative-control siRNA (20 μM each) using the Lipofectamine RNAiMAX transfection reagent (Thermo Fisher Scientific). Explants were cultured on nucleopore filters at 37 °C in a 5% CO2 atmosphere, in a Trowell-type organ culture containing a minimum essential medium with 10% KnockOut Serum Replacement formulation (Invitrogen), with the medium being changed every 2 days. After 2 days, some explants were homogenized using the TRizol reagent (Invitrogen), and cDNA from these samples was serially diluted for qRT-PCR to confirm target knockdown. After 10 days, the remaining explants were fixed in 4% paraformaldehyde, embedded in paraffin, serially sectioned (7 μm), and processed for H&E staining.

Renal capsule assay. E10 Runx2−/− and wild-type mouse mandibles were dissected under a stereomicroscope, and explants were transplanted beneath the kidney capsules of nude mice (KSN/Slc) together with the cationized gelatin sheet. Cationized gelatin (E7) was prepared, as described previously35,39,40. Briefly, cationized gelatin solution was freeze-dried to prepare cationized gelatin sheets, which were then cross-linked at 160 °C for 24 h. According to electronic scale measurements, each cross-linked cationized gelatin sheet weighed 1 mg. To impregnate phosphate-buffered saline (PBS) or Stealth siRNA into the cationized gelatin sheet, PBS or an aqueous solution containing Stealth siRNA (10 μL) was added to 1 mg of the cross-linked cationized gelatin sheet and then incubated at 37 °C for 1 h. The concentration of Stealth siRNA was adjusted with PBS to 322 μg/mL. Subcutaneous implantation was performed using a pair of fine tweezers and a microsurgery scalpel under a stereomicroscope. At 19 days post-transplantation, the mice were euthanized, and their kidneys were collected. The tissue attached to the kidney was removed and fixed in 4% paraformaldehyde, embedded in paraffin, and serially sectioned (7 μm), and processed for H&E staining. Some tissues were homogenized using the TRizol reagent (Invitrogen), and cDNA from these samples was subject to qRT-PCR to evaluate amelogenin and ameloblastin mRNA levels.

Micro-CT analysis. We performed 3D micro-CT scans (inspeXio SMX-100CT; Shimadzu, Kyoto, Japan) on the kidneys of mice at 19 days post-renal capsule transplantation. We converted CB files [512 x 512 pixels, 8 bits; voxel size, x:y:z = 1:1:1 (~ 0.06 mm/side)] to TIFF files, and 3D images were reconstructed and analyzed using computer imaging software (VGSTUDIO MAX 3.2; Volume Graphics GmbH., Heidelberg, Germany). Additionally, 3D-VR image of H&E-stained sections were reconstructed and analyzed using computer imaging software (Avizo 2019.2; Thermo Fisher Scientific, Waltham, MA, USA).

Immunohistochemistry. Examination of Stealth siRNA penetration into E10 wild-type mouse mandibular kidney capsules. E10 wild-type mouse mandibles were dissected, and explants were transplanted beneath the kidney capsule with gelatin sheets containing PBS or Alexa Fluor 488 (Thermo Fisher Scientific). At days 3, 6, and 10 post-transplantation, the mice were euthanized, and the tissue attached to the kidney was removed and fixed in 4% paraformaldehyde, embedded in paraffin, and serially sectioned (7 μm). Paraffin-embedded sections were subjected to immunostaining with primary anti-Alex Fluor 488 rabbit IgG Fraction (1:500) (A11094; Thermo Fisher Scientific, Waltham, MA, USA), and secondary goat and mouse anti-rabbit antibodies (Nichirei...
Bioscience, Tokyo, Japan). The sections were then counterstained with hematoxylin and dehydrated in a graded series of ethanol and xylene, after which coverslips were applied and the sections were viewed under a microscope.

Immuno- and histochimical assessment of amelogenin in Runx2-KO mouse mandibular kidney capsules. E10 Runx2+/− mice mandibles were dissected, and the explants were transplanted beneath the kidney capsule together with gelatin sheets containing Stealth RNAi siRNA #304. At day 19 post-transplantation, the mice were euthanized, and the tissue attached to the kidney was removed and fixed in 4% paraformaldehyde, embedded in paraffin, and serially sectioned (7 μm). The sections were immunostained with a primary anti-amelogenin antibody (1:500) (Hokudo, Sapporo, Japan), and secondary goat and mouse anti-rabbit antibodies (Nichirei Bioscience). The sections were then counterstained with hematoxylin and dehydrated in a graded series of ethanol and xylene, after which coverslips were applied and the samples were viewed under a microscope.

Statistical analysis. We presented categorical data as frequency and percentage. Statistical analyses were performed using the Wilcoxon rank-sum test. In animal experiments, it was not feasible to make an assumption of sample sizes statistically because we have never observed the development of cationized gelatin hydrogel as a DDS for Stealth siRNA during renal capsule transplantation of mouse mandibles. We did not perform any randomization. The investigators were not blinded for group allocations. Statistical analyses were conducted using SAS version 9.4 (SAS Institute Inc. Cary, NC).

Data availability

All data generated or analyzed during this study are included in this published article (and its Supplemental information files).

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**Author contributions**

K.T., Y.T1. Y.T2. and M.S. designed the research plan. S.M. preformed all the experiments. Analysis and interpretation of data were performed by H.B., H.K., A.MS., J.J., R.U., Y.N., T.K., H.H., and K.B., S.M. and K.T. wrote the main manuscript text. S.M. a prepared all figures. All authors reviewed and approved the manuscript.

**Competing interests**

The authors declare no competing interests.

**Additional information**

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