The Enamel Phenotype in Homozygous Fam83h Truncation Mice

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Abstract
Background: Truncation FAM83H mutations cause human autosomal dominant hypocalcified amelogenesis imperfecta (ADHCAI), an inherited disorder characterized by severe hardness defects in dental enamel. No enamel defects were observed in Fam83h null mice suggesting that Fam83h truncation mice would better replicate human mutations.

Methods: We generated and characterized a mouse model (Fam83hTr/Tr) expressing a truncated FAM83H protein (amino acids 1–296), which recapitulated the ADHCAI‐causing human FAM83H p.Tyr297* mutation.

Results: Day 14 and 7‐week Fam83hTr/Tr molars exhibited rough enamel surfaces and slender cusps resulting from hypoplastic enamel defects. The lateral third of the Fam83hTr/Tr incisor enamel layer was thinner, with surface roughness and altered enamel rod orientation, suggesting disturbed enamel matrix secretion. Regular electron density in mandibular incisor enamel indicated normal enamel maturation. Only mildly increased posteruption attrition of Fam83hTr/Tr molar enamel was observed at 7‐weeks. Histologically, the Fam83hTr/Tr enamel organ, including ameloblasts, and enamel matrices at sequential stages of amelogenesis exhibited comparable morphology without overt abnormalities, except irregular and less evident ameloblast Tomes’ processes in specific areas.

Conclusions: Considering Fam83h-/- mice showed no enamel phenotype, while Fam83hTr/Tr (p.Tyr297*) mice displayed obvious enamel malformations, we conclude that FAM83H truncation mutations causing ADHCAI in humans disturb amelogenesis through a neomorphic mechanism, rather than haploinsufficiency.

KEYWORDS
amelogenesis imperfecta, hair defects, knockout mouse, neomorphic mechanism, skin defects, truncation mutation
1 | INTRODUCTION

FAM83H (family with sequence similarity 83 member H on chromosome 8q24.3; OMIM *611,927) is a gene that was originally identified by gene prediction of the human genome through computational biology (Marchler-Bauer et al., 2005). Based upon the predicted 1,179 amino acids of human FAM83H protein, the only functional domain motif was the N-terminal domain (DUF1669) of unknown function that characterizes all eight members of the FAM83 protein family (Finn et al., 2016). Little attention was paid to FAM83H until it was reported to cause autosomal dominant hypocalciﬁed amelogenesis imperfecta (ADHCAI, OMIM #130900) (Kim et al., 2008). ADHCAI is a specific type of inherited dental enamel malformation in humans. The defective enamel is hypomineralized with a marked decrease in hardness and is lost following tooth eruption, which imposes esthetic and functional burdens on affected individuals (Wright, Deaton, Hall, & Yamauchi, 1995). However, in spite of the high prevalence of ADHCAI in North America and the many FAM83H mutations that have been identiﬁed in ADHCAI patients, the pathogenesis of this disease is largely unknown.

To date, 27 different FAM83H autosomal dominant truncation mutations have been associated with ADHCAI (Chan et al., 2011; Ding et al., 2009; El-Sayed, Shore, Parry, Inglehearn, & Mighell, 2010; Hart et al., 2009; Haubek et al., 2011; Hyun et al., 2009; Kantaputra, Intachai, & Auychai, 2016; Kim et al., 2008; Lee et al., 2008, 2011; Prasad et al., 2016; Song, Wang, Zhang, Yang, & Bian, 2012; Wang, Hu, Yang, Smith, Richardson, et al., 2015; Wright et al., 2009, 2011; Xin, Wenjun, Man, & Yuming, 2017; Zhang, Song, & Bian, 2015). Seven of these mutations have been found in multiple families. A contribution to the genetic etiology of ADHCAI has also been proposed for two heterozygous missense mutations, p.Ser342Thr (Pourhashemi et al., 2014) and p.Gly557Cys (Urzuza et al., 2015), but their role in ADHCAI causality should be considered with caution. According to the ExAC database, the p.Gly557Cys mutation has a minor allele frequency of 3.57% in the non-Finnish European population, suggesting that this mutation is most likely a neutral sequence variant, considering the rarity of ADHCAI. The p.Ser342Thr mutation has not been documented in the database, but the substituted amino acid is not highly conserved evolutionarily, and serine to threonine is usually not structurally significant and is predicted to be “benign” using the PolyPhen-2 prediction algorithm of functionality, suggesting that this mutation is also not likely to be disease-causing (Adzhubei, Jordan, & Sunyaev, 2013). Disregarding these two missense mutations of questionable causality, all of the other disease-causing FAM83H mutations are either nonsense or frameshift mutations that generate a premature termination codon in the most 3-prime exon of FAM83H. The resulting FAM83H transcripts apparently do not undergo nonsense mediated decay and are translated into truncated FAM83H proteins that include only the N-terminal region of FAM83H, extending no less than 287 amino acids and no longer than 694 amino acids (out of 1,179 amino acids in the full-length FAM83H protein). Neither truncation mutations outside of this speciﬁc region (287–694) nor other types of loss-of-function mutations seem to be disease-causing. For this evident mutational homogeneity, it has been suspected that a dominant negative effect or a gain of function, rather than haploinsufﬁciency, underlies the pathological mechanism of ADHCAI. We previously generated and characterized Fam83h null mice, further demonstrating this hypothesis (Wang, Hu, Yang, Smith, Richardson, et al., 2015). The lack of evident enamel defects in Fam83h heterozygous or null mice suggested that FAM83H is not required for normal enamel development in mice. A simple decrease of FAM83H function, including haploinsufﬁciency and dominant negativity, is not the cause of defective enamel in human ADHCAI. Instead, the truncated FAM83H from the mutant allele appears to impair normal enamel development by executing certain “toxic” effects (neomorphism) in ameloblasts, which cause the observed dominant enamel malformations.

In order to discern the “toxic” effects of truncated FAM83H in ameloblasts and unravel the pathogenesis of human ADHCAI, we generated a mouse model (Fam83hTr/Tr) by knocking in a truncation mutant Fam83h allele expressing only the N-terminal 296 amino acids of the FAM83H protein to recapitulate the human FAM83H p.Tyr297* mutation in mice. By characterizing the enamel phenotypes of Fam83hTr/Tr mice, we hypothesized potential disease mechanisms of ADHCAI at the cellular level and discussed difﬁculties and limitations of establishing a disease animal model to recapitulate a human disorder caused by neomorphic mutations.

2 | MATERIALS AND METHODS

2.1 | Generation of knockin mice (Fam83hTr)

The knockin construct was designed and the mice (species: Mus musculus, strain C57BL/6) were generated by Ozgene (Perth, Australia). The Fam83hTr mouse was generated by replacing Exon 4 of the endogenous Fam83h with a mini-cDNA that introduced an early translation termination codon. The mini-cDNA was composed of the whole of Exon 4 of Fam83h replacing Exon 4 of the endogenous FAM83H, which is transcribed using its native promoter and translated from its native initiation codon, but generates a transcript encoding an open reading frame containing only the first 296 codons of wild-type
Fam83h, followed by multiple polyadenylation signals. This mutant transcript is predicted to produce a truncated mouse Fam83H protein (normally 1,209 amino acids) with only the N-terminal 296 amino acids (FAM83H1-296) within the natural physiological context of Fam83h expression in vivo. This mouse model mimics a specific human FAM83H disease-causing mutation, p.Tyr297*. The deleted Fam83h and inserted NLS-lacZ sequences are provided in Figures S3–S4.

2.2 | Physical assessment and photography

Heterozygous mutant (Fam83h+/Tr) mice were mated to generate three mouse Fam83h genotypes (+/+; +/Tr, and Tr/Tr), which were then evaluated for their physical appearance, activity, growth rate, size difference, food intake, and fertility. The homozygous (Fam83hTr/Tr) mice were viable and fertile. For gross evaluation of the dental enamel, mice were put under anesthesia using isoflurane, and their incisors were inspected under a dissection microscope. Seven-week-old mice were sacrificed and fixative-perfused with 4% paraformaldehyde (PFA). The mandibles were removed and sliced through the mental symphysis with a razor blade to generate hemimandibles. These were carefully dissected free of soft tissues under a stereoscopic microscope using tissue forceps and a spoon excavator. The hemimandibles were submerged in 1% NaClO for 5 min, rinsed, air dried, and photographed using a Nikon digital camera DXM1200 (Melville, NY).

2.3 | Molar histology

For histology of developing molars, Day 5, 11, and 14 mouse heads were quickly dissected free of skin, cut in half, and immersed in 4% PFA fixative overnight at 4°C and decalcified at 4°C by immersion in 1 L of 4.13% disodium EDTA (pH 7.3) with agitation. The EDTA solution was changed every other day for 30 days. The samples were washed in PBS at 4°C 4–5 times every 0.5–1 hr, washed overnight, postfixed for 2 hr in 1% osmium tetroxide/1.5% potassium ferrocyanide, dehydrated using an acetone gradient, embedded in Epon812 substitute, semi thin-sectioned and stained with 0.1% toluidine blue as described elsewhere (Smith, Hu, et al., 2011). Three maxillary (one from each genotype), two Fam83h+/+, two Fam83h+/Tr, and five Fam83hTr/Tr mandibular incisors were processed for longitudinal sectioning. Three Fam83h+/+ and four Fam83hTr/Tr mandibular incisors were processed for cross sectioning at 1 mm increments as described previously (Hu et al., 2016).

2.5 | Sodium dodecyl sulfate polyacrylamide gel electrophoresis (SDS-PAGE)

The detailed protocol of protein extraction from mouse molars and the analyses were previously described (Wang, Hu, Yang, Smith, Nunez, et al., 2015; Yamakoshi et al., 2011). Briefly, mouse maxillary and mandibular first Fam83h+/+; Fam83h+/Tr; Fam83hTr/Tr; Fam83h−/− molars were extracted from mouse pups at D5, D11, and D14 using a dissecting microscope, stripped of soft tissue, and the mineralized portion of the teeth was incubated in 1 ml of 0.17 N HCl/0.95% formic acid for 2 hr at 4°C. After the undissolved materials were removed, the crude protein extract was buffer exchanged with 0.01% formic acid. The concentrate containing proteins extracted separately from the four molar genotypes was raised back to 250 μl of 0.01% formic acid and used for subsequent analysis by SDS-PAGE stained with Coomassie Brilliant Blue (CBB). The amount of protein applied per lane for SDS-PAGE was ~1/6 of the extract from a single molar. Recombinant mouse amelogenin (rM179) was used as a size control (Simmel et al., 1994).

2.6 | Backscattered scanning electron microscopy

The procedures of backscattered scanning electron microscopy (bSEM) were described previously (Smith, Richardson, et al., 2011). Soft tissue was removed from the left and right hemimandibles of 7-week-old mice, and cross-sectioned at 1 mm increments from the basal (growing) end of the incisors and imaged by bSEM using a JOEL 7,800 SEM at 20 kV (JOEL USA, Inc; Peabody, MA, USA). For whole incisor surface imaging, the bony soft tissue and bony caps covering the mandibular incisors were carefully removed and examined at 50X magnification with a Hitachi S-3000N variable pressure SEM using the backscatter mode at 25 kV and 20 pascal pressure.
For molar surface SEM, the molars were prepared as follows: The D14 hemimandibles were submerged in 4% PFA overnight, carefully dissected of soft tissues under a stereoscopic microscope, submerged in 1% NaClO for 20 min, rinsed, and acetone dehydrated (30%, 50%, 70%, 80%, 90%, 100%). The hemimandibles were mounted on metallic stubs using conductive carbon cement and carbon coated to increase conductivity and examined using a Hitachi (Century City, Los Angeles, CA) S-3000N variable pressure SEM using the backscatter mode.

### 2.7 Hardness evaluations

The procedures for tissue preparation and microhardness testing were previously described (Wang, Hu, Yang, Smith, Nunez, et al., 2015). In brief, hemimandibles from 7-week-old mice were cleaned free of soft tissue and embedded in Epon resin following graded acetone dehydration. After polymerization at 60°C, the incisors were cross-sectioned 8 mm from the apical end of the incisor, approximately at the level of the crest of the alveolar bone close before the incisor erupts into the mouth. The cross-sectioned incisor was reembedded in Castolite AC (Eager Polymers, Chicago, IL) using 25-mm SteriForm molds (Struers Inc., Westlake, OH), allowed to harden overnight, and polished.

Microhardness testing was performed using a LM247AT microhardness tester (Leco Corp., St. Joseph, MI) with a load of 25 g for 10 s with a Knoop tip to obtain a Knoop hardness number (KHN). Measurements were made at 500X magnification. Indentations were placed in the outer, middle, and inner enamel as well as the dentin as a control reading for a total of four indentations per row. This series was performed three times in each animal, for a total of twelve points per animal. Hardness Data points were treated as independent, unweighted numbers and subjected to one-way ANOVA than Tukey’s HSD test using calculation spreadsheet at http://vassarstats.net/anova1u.html. Statistical significance was determined at \( p < 0.01 \).

Nano hardness test was done on (Hysitron TI 950 Triboindenter, Hysitron-Bruker, Minneapolis, MN, USA). Using the nano DMA indenter, the loading force was 8,000 µN, 15 indents on each incisor spread among dentin, inner enamel, middle enamel and outer enamel. The data was analyzed by using Hysitron Troboscans.

### 2.8 Microcomputed tomography (µCT)

Sample preparation and imaging process for microcomputed tomography (µCT) were conducted as previously described (Yang et al., 2015). Briefly, the agarose-embedded hemimandibles were scanned at 8 µm and analyzed with the SCANCO µCT-100 series microcomputed tomography system. The data were analyzed using the MiiL program (Medical Image Illustrator; National Applied Research Laboratories, Taipei, Taiwan). For calculating the hard tissue volumes from the 3D molar images, the contour of each first molar was outlined by marking the border of the tooth on each scanning section. Three dimensional images were generated within the contour defined area. Based upon the distinct radiodensity of each tissue, the enamel, dentin, and pulp of the molars were isolated using density threshold ranges to measure the tissue volumes. The threshold range for measuring the pulp was 0 to 136, for dentin was 137 to 225, and for enamel was 226 to 255. Pulp, dentin, and enamel volumes as defined by these threshold ranges were calculated using the MiiL Program. The total volume of the whole tooth was the sum of volumes from its enamel, dentin, and pulp. The proportion of each tissue within a tooth was then calculated. The volume ratios of enamel to dentin (E/D) and enamel to dentin-pulp complex (E/D + P) were also evaluated.

## 3 RESULTS

### 3.1 Generation of truncated Fam83h-knockin mice

In order to recapitulate human FAM83H disease-causing mutations and investigate the pathogenesis of human ADHCA1, we generated a mouse model expressing a truncated (Tr) FAM83H protein in the C57BL/6-background. The structures of wild-typeFam83h, the Fam83h targeting construct, the Cre-inverted intermediate, and the final Fam83hTr gene and strategy for PCR genotyping are provided in Figure S1. We also provide relevant DNA sequences, such as the wild-type mouse Fam83h gene sequence and structure (Figure S2), the knockin intermediate following removal of the selection genes using Flp-FRT recombination (Figure S3), and the final Fam83hTr knockin sequence (Figure S4).

Intercrosses of heterozygous (Fam83hTr/+) mice produced pups of three genotypes at the expected Mendelian ratio. The homozygous (Fam83hTr/Tr) mice were viable and fertile. At birth, they were indistinguishable from their wild-type (Fam83h+/+) and heterozygous (Fam83hTr/+) littermates. However, when inspected at 7-weeks, the homozygous mutants (Fam83hTr/Tr) appeared normal in size but exhibited a sparse and scruffy coat, which allowed them to be readily discerned from their heterozygous and wild-type littermates. Despite the coat distinctness of 7-week Fam83hTr/+, their incisors, when inspected under a dissection microscope (Figure 1), appeared normal with glossy enamel surfaces and sharp incisal tips, comparable to those of their Fam83hTr/+ and Fam83h+/- littermates. However, when carefully viewed from the lateral perspective (Figure 1b), the incisor enamel did not show a normal color of glazy yellow and a continuous curved line of light reflection, but instead exhibited a dull-white appearance with disrupted patchy areas of light reflection. Basic tooth

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[Note: The text continues with additional details related to experiments, results, and conclusions pertinent to the research described.]
morphology looked comparable among the three genotypes, although the molar enamel surface textures were not as smooth as those of the wild-type mice (Figure 1c). Notably the extensive posteruption attrition of enamel that is pathognomonic of the ADHCAI phenotype caused by heterozygous FAM83H truncation mutations in humans was absent from the mouse heterozygous molars and mild even in the Fam83hTr/Tr molars.

3.2 | Enamel defects in Fam83hTr/Tr molars

To assess the potential enamel phenotypes in detail, we examined 7-week molars using backscattered scanning electron microscopy (bSEM; Figure 2; Figure S5). The basic morphology of the molars was comparable in the three genotypes, although the Fam83hTr/Tr molars sometimes appeared to be mildly smaller than the Fam83h+/+ and Fam83h+/Tr molars, and their cusps seemed slender compared to those of the Fam83h+/+ and Fam83h+/Tr mice, suggesting that their enamel layer might be thinner. The Fam83hTr/Tr molar surfaces were less smooth than those of the Fam83h+/+ and Fam83h+/Tr molars, confirming the impression made under dissection microscope. When inspected from the occlusal view, the cusp tips of Fam83hTr/Tr first molars exhibited only slightly larger wear facets, suggesting that the Fam83hTr/Tr molars had undergone similar mild attrition relative to the Fam83h+/+ and Fam83h+/+ molars.

We removed the overlying soft tissue covering the unerupted (but nearly erupted) Day 14 molars and characterized them using bSEM (Figure 3). The unerupted Fam83hTr/Tr molars were similar to the 7-week Fam83hTr/Tr molars that had functioned in occlusion for 5 weeks by also exhibiting enamel surface roughness and cusp slenderness, indicating that these phenotypes resulted from developmental defects rather than posteruption changes.

To better assess the possibilities of smaller molar size and thinner enamel suggested by the dissection and bSEM microscopy, we performed µCT analyses to measure the volumes of enamel, dentin, and pulp in mandibular first molars at Day 14. Specific ranges of µCT threshold values were set to isolate each tissue (pulp, dentin and enamel): 0–136 for pulp, 137–225 for dentin, and 226–255 for enamel. The means of the measurements were calculated from six wild-type, five heterozygous, and six homozygous D14 mice. There were no statistically significant differences in the volumes of the three dental tissues between the wild-type and heterozygous mice. In contrast, the total molar volume, and the volumes of dentin, enamel were reduced between the Fam83h+/+ and
the Fam83h<sup>Tr/Tr</sup> mice, and these differences were statistically significant (Figure 4). The total tooth, dentin, and enamel volumes decreased to 84%, 87%, 62% of those of the wild-type, respectively. Overall, the mandibular first molars of the Fam83h<sup>Tr/Tr</sup> mice were smaller than those of the wild-type. As the proportion of dentin increased in the homozygous (77%) relative to the wild-type (75%), while the proportion of enamel fell in the homozygous (9.7%) relative to the wild-type (13.2%), it is evident that the Fam83h<sup>Tr/Tr</sup> molars are smaller than those of the wild-type principally because of a reduction in enamel volume, confirming a hypoplastic enamel defect in Fam83h<sup>Tr/Tr</sup> molars.

We analyzed the histology of developing maxillary first molars at postnatal D5, D11, and D14 for three genotypes (Figure 5; Figure S6). All D5 first molars showed typical tall-columnar secretory stage ameloblasts with accumulation of eosinophilic extracellular matrices in the enamel. Similarly, in D11 first molars of all genotypes, the maturation stage ameloblasts appeared comparable adjacent to an enamel space with some residual matrix. At D14, when the
molar was about to erupt into the oral cavity, all molars of the three genotypes exhibited a thin layer of reduced enamel epithelium outlining the enamel space containing little residual matrix. Odontoblasts, developing dentin-pulp complexes, developing roots, and surrounding periodontal tissues of D5, D11, and D14 first molars seemed to be comparatively normal in Fam83h+/Tr and Fam83hTr/Tr mice. Light microscope histology of developing first molars did not provide evidence of pathology that could explain the enamel hypoplasia or surface roughness.

We also utilized SDS-PAGE to analyze crude protein extracts from D5, D11, and D14 first molars of Fam83h+/+, Fam83h+/Tr, and Fam83hTr/Tr mice (Figure S7). No clear differences in the amount of matrix proteins were observed among the first molars of the different genotypes at D5, D11, and D14, which was consistent with the molar histology findings that ameloblast form and enamel architecture were not grossly altered in the Fam83h+/Tr and Fam83hTr/Tr mice. Light microscope histology of developing first molars did not provide evidence of pathology that could explain the enamel hypoplasia or surface roughness.

FIGURE 4 MicroCT Analyses of D14 Mandibular Molars. Day 14 precedes slightly the eruption of the first molar into function, before the potential effects of attrition on the volumes of enamel and dentin can occur. No statistically significant molar volumes were observed between the Fam83h+/+ and Fam83h+/Tr and are not displayed. Total molar volume was reduced in the Fam83hTr/Tr to 84% that of the wild-type. The volume of dentin was reduced by a lesser amount (to 87%), indicating that part of the total volume reduction was due to an overall reduction in tooth size. The volume of enamel was reduced in the Fam83hTr/Tr to 62% that of the wild-type indicating that most of the reduction in molar size was due to a reduction in enamel volume.

3.3 | Enamel defects in Fam83hTr/Tr mandibular incisors

The mineralization of mandibular incisors was analyzed by comparing backscattered scanning electron microscopy (bSEM) images of 1-mm incremental cross sections (from level 2 to 8) progressing from early (apical) to late (incisal) stage of development along the continuously growing mandibular incisors from Fam83h+/+, Fam83h+/Tr, and Fam83hTr/Tr mice. The incisor cross sections from all levels were imaged at three magnifications and the three genotypes compared at each magnification. These magnifications (from low to high) show the entire incisor cross sections (Figure S8), the enamel layer and underlying dentin (Figure S9), and the enamel layer at the height of contour (Figure S10), where the enamel is thickest and its rod pattern most evident. These comparisons, along with the study of bSEM image panels showing six Fam83h+/+, six homozygous (Fam83hTr/Tr) (Figures S11–S16), and one wild-type (Fam83h+/+) (Figure S23) incisors at 1 mm cross sections at all three magnifications, demonstrated virtually normal progression of dentin and enamel mineralization in all three genotypes. The gist of these findings is evident in Figure 6, which shows a panel of bSEM images of incisor cross sections at level 8 that show the enamel layer after it had completed maturation and was about to erupt into the oral cavity. The enamel layer at level 8 in the WT, heterozygous (Fam83h+/+) and homozygous (Fam83hTr/Tr) mice exhibited comparable thickness (from the DEJ to the enamel
surface at the height of contour), electron density (reflecting degree of mineralization) and overall enamel rod patterns at sequential sections among the three genotypes, indicating that expression of the truncated FAM83H protein did not interfere with enamel and dentin mineralization, making it unlikely that enamel matrix secretion was disturbed. However, when inspected carefully, the lateral (lateral) 1/3 of the enamel layer of \( \text{Fam83h}^{\text{Tr/Tr}} \) incisors was thinner relative to the \( \text{Fam83h}^{+/+} \) and \( \text{Fam83h}^{+/+} \) incisors (Figure 7). While the labial enamel surface of a normal incisor exhibited a smooth line gradually approaching the lateral cervical margin on a cross-section, the contour of \( \text{Fam83h}^{\text{Tr/Tr}} \) enamel abruptly steepened down at the lateral 1/3. This was evident at incremental sections starting from level 4, which represented the postsecretory (maturation) stage of enamel formation. In contrast, the dentin and surrounding alveolar bone of mandibular incisors in all three genotypes appeared to be comparable on bSEM cross sections.

To quantify this hypoplastic enamel defect, we analyzed the mesial and lateral enamel areas of level 8 cross sections from 10 \( \text{Fam83h}^{+/+} \) and eight \( \text{Fam83h}^{\text{Tr/Tr}} \) 7-week mandibular incisors (Figure 8; Figure S24). A bisecting line perpendicular to a line segment connecting the mesial cervical margin and the lateral cervical margin was used to divide the incisor enamel into mesial and lateral parts, and the areas of each part and the total enamel were calculated. In the wild-type incisor, the average area of total enamel was around 73,700 µm², roughly three-fifths (45,600 µm²) of which was mesial enamel and two-fifths (28,100 µm²) was lateral enamel. In contrast, the \( \text{Fam83h}^{\text{Tr/Tr}} \) incisor exhibited significantly smaller areas of total and lateral enamel (66,800 and 22,300 µm² respectively), but not mesial enamel (44,500 µm²). While the mesial enamel area appeared to be comparable between the wild-type and \( \text{Fam83h}^{\text{Tr/Tr}} \) incisors, the lateral enamel area of the \( \text{Fam83h}^{\text{Tr/Tr}} \) incisor was less than 80% that of the wild-type incisor. This uneven reduction of enamel area changed the proportion of lateral to mesial enamel from 2:3 to 1:2, confirming a thickness defect in enamel on the lateral aspect of a \( \text{Fam83h}^{\text{Tr/Tr}} \) mandibular incisor.

To assess potential alteration of enamel functionality in \( \text{Fam83h}^{\text{Tr/Tr}} \) mice, we conducted Knoop hardness testing (KHT) and Nano hardness testing (NHT) of dentin, the inner enamel (near dentin), the middle enamel, and outer enamel
FIGURE 6  bSEM of 7-week mandibular incisor cross sections. These incisor cross sections show the enamel layer at a level near the crest of the alveolar bone (level 8), which is prior to eruption into the oral cavity but after the enamel has undergone extensive maturation. The enamel looks similar in the WT (+/+), heterozygous (+/Tr) and homozygous (Tr/Tr) mice. The enamel of the Fam83hTr/Tr incisors appears to have normal centro‐labial thickness, rod patterns and degree of mineralization. The only difference is in the contour of the enamel as it approaches the lateral side, so that the enamel layer is thinner than normal as it approaches the lateral cervical margin.

3.4  |  Histological analyses of Fam83h incisors at 7-weeks

We analyzed the histology of longitudinal sections of continuously growing maxillary (Figures S28–S30) and mandibular (Figures S31–S48) incisors of 7-week Fam83h+/+, Fam83h+/Tr, and Fam83hTr/Tr mice, which exhibited all stages of enamel formation. The basic morphology of the enamel organ, including ameloblasts, and enamel matrices at differentiation, secretory, transition, and maturation stages appeared comparable in the three genotypes, although the secretory stage ameloblasts of Fam83hTr/Tr incisor gave the impression of being shorter, with a likely increased nucleus-to-cell height.
ratio, compared to those of Fam83h\(^{+/+}\) and Fam83h\(^{+/Tr}\) incisors (Figure 10). The Fam83h\(^{Tr/Tr}\) incisor also exhibited normal onset of ameloblast differentiation and transition through all stages of amelogenesis without apparent delay, advance, or interruption. Furthermore, a clear enamel space with no residual staining was consistently observed beneath late maturation stage ameloblasts (Figures S28, S32, S34, S36, S38 and S40), which indicated that the process of enamel maturation in the Fam83h\(^{Tr/Tr}\) incisor had proceeded normally.

Being aware of a hypoplastic defect at the lateral aspect of Fam83h\(^{Tr/Tr}\) incisor, we examined the histology of amelogenesis in 1-mm incremental cross sections (from level 1 to 10; Figures S49–S56) analogous to the previously-mentioned bSEM analyses, including high magnification montages of the incisor cross sections (Figures S57–S61). Ameloblasts reverse their secretory polarity during differentiation to allow for extracellular release of large amounts of proteins from plasma membrane surfaces that were originally the embryonic bases of the cells (Smith & Nanci, 1995). In general, level 1 showed the onset of secretory stage with elongated ameloblasts being reversely-polarized and developing Tomes' processes, while the midsecretory stage could be observed at level 2 showing tall-columnar ameloblasts with well-established Tomes' processes tilting mesially or laterally. The transition stage usually appeared around level 3, followed by the maturation stage extending from levels 3 through 8. With gradual removal of enamel matrix proteins and increased mineralization throughout the maturation stage, staining of the enamel matrix was barely observed after level 5. Consistent with the findings from longitudinal sections, the Fam83h\(^{Tr/Tr}\) incisors exhibited comparable histology without overt
morphological abnormalities at all section levels, relative to the Fam83h\textsuperscript{+/+} incisors. However, at level 2 (mid-secretory stage), the ameloblasts over the lateral aspect of Fam83h\textsuperscript{Tr/Tr} incisor showed less evident Tomes' processes than those of wild-type at the comparable area (Figure 11). Also, in some Fam83h\textsuperscript{Tr/Tr} samples, the Tomes' processes of ameloblasts at the central area of level 2 were irregular in the enamel layer, which was rarely observed in the wild-type samples (Compare the wild-type Level 2 ameloblasts near the lateral cervical margin of Figure S57 to the Fam83h\textsuperscript{Tr/Tr} ameloblasts in the same position in Figures S58–S61).

4 | DISCUSSION

We recently demonstrated that Fam83h null mice exhibited no overt enamel defects, which suggested a nonessential role for FAM83H in enamel formation and supported a
neomorphic disease mechanism of human autosomal dominant hypocalcified amelogenesis imperfecta (ADHCAI) (Wang, Hu, Yang, Smith, Richardson, et al., 2015). An amorphic (loss of function) disease mechanism for ADHCAI is ruled out in humans because only truncation mutations in a specific region of the last exon cause ADHCAI, whereas missense mutations in conserved positions and nonsense mutations in other parts of the gene that could cause a loss of function are not observed in ADHCAI patients. In mice, an amorphic mechanism for ADHCAI is ruled out because the Fam83h heterozygous and null mice do not exhibit an enamel phenotype (Wang, Hu, Yang, Smith, Richardson, et al., 2015). In this study we generated knockin (Fam83hTr/Tr) mice with a truncation defect (p.Tyr297*) homologous to those that cause human ADHCAI. The (Fam83hTr/Tr) mouse molars showed rough enamel surfaces and slender cusps while mandibular incisors showed enamel hypoplasia in their lateral enamel, indicating disturbed matrix secretion during enamel formation.

The mainly hypoplastic phenotype of Fam83hTr/Tr enamel indicated that the secretory stage was more affected by the truncated FAM83H protein than was the maturation stage. We previously showed that in mouse molars, Fam83h expression in ameloblasts was generally weak and barely detectable until the transition to the maturation stage (Wang, Hu, Yang, Smith, Richardson, et al., 2015). This inconsistency suggests that mouse secretory stage ameloblasts are more highly vulnerable to the “toxic” effect of truncated FAM83H, given that their expression of the protein is low. Maturation stage ameloblasts were more resistant, since the molar enamel hardness of Fam83hTr/Tr mice was only slightly reduced. Although Fam83hTr/Tr enamel was malformed, the defects were considerably different from those observed in ADHCAI patients. In human ADHCAI, the enamel of affected individuals is of normal thickness in unerupted or newly erupted teeth, but is hypomineralized, soft, and undergoes rapid attrition following eruption. It is unknown if human ameloblasts express FAM83H at higher levels than mouse ameloblasts, which might account for the more severe enamel defects in ADHCAI patients. Alternatively, human ameloblasts, particularly the maturation stage ones, might be more sensitive to the “toxic” effect of the mutant FAM83H. This potential difference in FAM83H expression level and ameloblast susceptibility to the mutant protein might explain why enamel defects could be seen in homozygous (Fam83hTr/Tr), but not heterozygous (Fam83h+/Tr) mutant mice, while a single copy of mutant FAM83H can cause ADHCAI in humans.
An interesting feature of *Fam83h*^Tr/Tr^ incisors was that enamel defects were only evident on the lateral aspect. The mutant FAM83H protein in mice more significantly impacts appositional growth of enamel over the lateral aspect (of the incisor) than the mesial, causing a localized hypoplastic defect. In that area the enamel was thinner, but also the normal pattern of enamel rod rows was lost (Figure 7). Interestingly, the altered *Fam83h*^Tr/Tr^ incisor did not seem to affect enamel functionality, namely hardness, since we did not detect significant differences with either KHT or NHT.

Tomes’ processes are cell projections of secretory stage ameloblasts into the enamel matrix. They are considered to be critical for establishment of enamel rod-interrod structure. Aberration of Tomes’ processes on the lateral aspect of incisors might explain the hypoplastic enamel in this area of *Fam83h*^Tr/Tr^ incisors. Kuga *et al.* recently showed that FAM83H and its associated protein CK-1ε colocalized with keratin filaments in mouse ameloblasts, and demonstrated that FAM83H and CK-1 regulate the organization of the keratin cytoskeleton, and maintain the formation of desmosomes in human ameloblastoma cell lines (Kuga *et al.*, 2016). Formation of Tomes’ processes by ameloblasts reflects a significant reorganization of the cytoskeleton, so perhaps truncated FAM83H disturbs the organization of the cytoskeleton, although the normal FAM83H protein is not critical for proper organization of the keratin cytoskeleton in ameloblasts, given the fact that there are no evident enamel defects in *Fam83h* null mice (Wang, Hu, Yang, Smith, Richardson, *et al.*, 2015).

Another interesting finding of *Fam83h*^Tr/Tr^ enamel defects was that the molars appeared more affected and showed more prominent hypoplastic defects than the incisors did. This discrepancy might result from a fundamental difference in enamel structure between molars and incisors. The
spatial distribution and three-dimensional arrangement of enamel types, namely schmelzmuster, in rodents have been well characterized and documented by paleontologists for phylogenetic purposes (von Koenigswald, 2004a, 2004b; von Koenigswald & Clemens, 1992). In most myomorph rodents, including mice (Mus musculus), the enamel microstructure of incisors comprises an inner layer of uniserial lamellar enamel (inner enamel) and an outer layer of radial enamel (outer enamel), as described above (von Koenigswald & Clemens, 1992; Moinichen, Lyngstadaas, & Risnes, 1996). The inner layer exhibits a decussation structure on cross sections and an architecture of Hunter-Schreger Bands (HSB) on sagittal (longitudinal) sections. On the other hand, the molars showed a more complicated enamel microstructure, named C-type schmelzmuster (von Koenigswald, 2004b). A basal ring of lamellar enamel (BRLE) surrounds the cervical part of the crown, covered by an outer layer of radial enamel, while the upper part of the enamel cap is formed by mostly radial enamel with rods perpendicular to the enamel surface. The uniserial HSBs in incisors are identical to the BRLE in molars. The truncated FAM83H protein disturbed the ameloblast Tomes’ processes and altered the enamel rod profiles they produced. These effects were more significant on radial enamel with rods perpendicular to the enamel surface, which comprise the greater part of a molar crown, contributing to the surface roughness of Fam83hTr/Tr molars. In contrast, the incisor outer enamel with rods running parallel to the enamel surface might be less susceptible to the altered rod–interrod profile caused by truncated FAM83H and therefore exhibit a minor hypoplastic defect. Interestingly, mice with ablation of serotonin 2B receptor (Htr2b null mice) have a similar molar phenotype of generally thinner enamel with surface roughness, and the enamel microstructure and rod profiles of Htr2b−/− molars were also abnormal (Dimitrova-Nakov et al., 2014; Harichane et al., 2011). Although the alteration of the rod profiles of molar enamel was not well detailed, primarily due to the high complexity of molar enamel microstructure and a lack of standardized methodology for molar characterization, it was clearly indicated that enamel analysis of mutant genotypes based exclusively on incisor defects can be biased (Goldberg, Kellermann, Dimitrova-Nakov, Harichane, & Baudry, 2014). Correspondingly, the Fam83hTr/Tr mice in this study also demonstrated distinct enamel phenotypes between incisors and molars, indicating a necessity of comprehensive characterization of both tooth types in a given mouse model and an imperative of developing standardized methods to investigate molar enamel microstructure.

We previously showed that Fam83h null mice died at around 2 weeks, and the few older survivors were generally weak, suggesting an essential role of FAM83H in certain critical physiological functions (Wang, Hu, Yang, Smith, Richardson, et al., 2015). Nonetheless, in this study, Fam83hTr/Tr mice appeared to be viable and fertile. This discrepancy in viability suggested that the N-terminal which is composed of 296 amino acids of FAM83H is sufficient for its critical functions. In other words, the truncated part of the protein in the C-terminus, composed of about 900 amino acids, is not necessary for viability functions. The protein family of FAM83 is characterized by a highly conserved N-terminal domain of unknown function, DUF1669, comprising ~300 amino acids (Cipriano et al., 2014). Except for this domain, the eight members of FAM83 family have vastly different C-terminal regions with variable lengths. The FAM83 proteins were recently identified as novel transforming oncogenes that function as intermediaries in EGFR/RAS signaling, and the oncogenic properties were dependent upon the conserved DUF1669 domain (Cipriano et al., 2012, 2014). It was also demonstrated that these proteins, including FAM83H, were overexpressed in many human tumors (Cipriano et al., 2014; Snijders et al., 2017). Therefore, FAM83H might play a role in cell proliferation or survival mediated by EGFR/RAS signaling, presumably through its DUF1669 domain, which is essential for certain critical physiological functions. In this regard, complete loss of FAM83H in Fam83h null mice would be fatal. In contrast, the Fam83hTr/Tr mice, although only expressing the first 296 amino acids of FAM83H containing the DUF1669 domain, would be able to survive. In other words, for certain cells other than ameloblasts, the mutant (truncated) FAM83H protein is not only not “toxic”, but can serve critical functions. Nevertheless, despite the viability, Fam83hTr/Tr mice exhibited sparse and scruffy coat as did Fam83h null mice (Wang, Hu, Yang, Smith, Richardson, et al., 2015), which suggested that the DUF1669 domain of FAM83H was not sufficient to serve its functions in hair follicle cells. Since FAM83H is believed to interact with keratin through its C-terminus (Kuga et al., 2013), FAM83H might be critical for maintaining the homeostasis of hair follicles via this interaction. Therefore, loss of FAM83H C-terminus, in both Fam83h null and Fam83hTr/Tr mice, leads to hair pathology.

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CONFLICTS OF INTEREST

The authors have no conflict of interest to declare regarding this manuscript.

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