STIM1 R304W in mice causes subgingival hair growth and an increased fraction of trabecular bone

Thilini H. Gamagea,1, Emma Lenglea,1, Gjermund Gunnesb, Helen Pullisaarc, Ashjorn Holmgreana, Janne E. Reselandd, Else Merckolle, Stefania Cortif,g, Masahiro Mizobuchih, Raul J. Moralesi, Leonidas Tsiokasj, Geir E. Tjønnfjordk, Rodrigo S. Lacruzl, Staale P. Lyngstadaasd, Doriana Misceoa,2, Eirik Frengen,a,⁎

a Department of Medical Genetics, Oslo University Hospital and University of Oslo, Oslo, Norway
b Faculty of Veterinary Medicine, Norwegian University of Life Sciences, Norway
c Department of Orthodontics, Institute of Clinical Dentistry, University of Oslo, Oslo, Norway
d Department of Biomaterials, Institute of Clinical Dentistry, University of Oslo, Oslo, Norway
e Department of Radiology and Nuclear Medicine, Oslo University Hospital, Oslo, Norway
f Neuroscience Section, Department of Pathophysiology and Transplantation, Dino Ferrari Centre, University of Milan, Milan, Italy
g Department of Neurology, Nakamura Memorial Hospital, Sapporo, Japan
h CHRU, Hôpital Gui de Chauliac, Montpellier, France
i Department of Cell Biology, University of Oklahoma Health Sciences Center, Oklahoma, USA
j Department of Haematology, Oslo University Hospital and Institute of Clinical Medicine, University of Oslo, Oslo, Norway
k Department of Basic Science and Craniofacial Biology, New York University College of Dentistry, New York, USA

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A B S T R A C T

Calcium signaling plays a central role in bone development and homeostasis. Store operated calcium entry (SOCE) is an important calcium influx pathway mediated by calcium release activated calcium (CRAC) channels in the plasma membrane. Stromal interaction molecule 1 (STIM1) is an endoplasmic reticulum calcium sensing protein important for SOCE.

We generated a mouse model expressing the STIM1 R304W mutation, causing Stormorken syndrome in humans. Stim1R304W/R304W mice showed perinatal lethality, and the only three animals that survived into adulthood presented with reduced growth, low body weight, and thoracic kyphosis. Radiographs revealed a reduced number of ribs in the Stim1R304W/R304W mice. Microcomputed tomography data revealed decreased cortical bone thickness and increased trabecular bone volume fraction in Stim1R304W/R304W mice, which had thinner and more compact bone compared to wild type mice. The Stim1R304W/+ mice showed an intermediate phenotype. Histological analyses showed that the Stim1R304W/+ mice had abnormal bone architecture, with markedly increased number of trabeculae and reduced bone marrow cavity. Homozygous mice showed STIM1 positive osteocytes and osteoblasts. These findings highlight the critical role of the gain-of-function (GoF) STIM1 R304W protein in skeletal development and homeostasis in mice. Furthermore, the novel feature of bilateral subgingival hair growth on the lower incisors in the Stim1R304W/R304W mice and 25% of the heterozygous mice indicate that the GoF STIM1 R304W protein also induces an abnormal epithelial cell fate.

1. Introduction

Calcium signaling plays an important role in bone development and homeostasis [1]. One important Ca²⁺ entry pathway in non-excitable cells is the store-operated Ca²⁺ entry (SOCE) [2,3]. SOCE is mediated by stromal interaction molecule 1 (STIM1), a single-pass transmembrane protein, which resides in the endoplasmic reticulum (ER) membrane and senses ER Ca²⁺ levels [4]. When ER Ca²⁺ stores are low,
STIM1 oligomerizes and translocates to ER-plasma membrane (ER-PM) junctions [5,6]. At these junctions, STIM1 interacts with ORAI1, a tetraspanning PM protein. ORAI1 forms the ion-conducting pore of the Ca\(^{2+}\) release-activated Ca\(^{2+}\) channel (CRAC) and enables Ca\(^{2+}\) influx across PM [2,3,7–9]. The study of patients and mouse models with mutations in ORAI1 or STIM1 genes contribute to the understanding of the function of these proteins through phenotypes related to dysregulated SOCE [8,10–15].

Mutations in the ORAI1 or STIM1 genes cause diseases in human [9–11,13–15]. Bi-allelic loss-of-function (LoF) mutations in STIM1 or ORAI1 result in absence of SOCE and patients present with Immunodefucence 10 (IMD10, OMIM #612783) or Immunodefucence 9 (IMD9, OMIM #612782) [12,15], respectively. The clinical presentation includes immunodeficiency, autoimmunity, ectodermal dysplasia, muscular hypotonia, and severe dental enamel defects, which highlights the role of CRAC in tooth development [8,9,12,15–17]. Heterozygous gain-of-function (GoF) mutations in STIM1 and ORAI1 in humans cause constitutive SOCE resulting in a spectrum of diseases ranging from isolated tubular aggregate myopathy (TAM) (TAM1, OMIM #160565, TAM2, OMIM #615883) to Stormorken syndrome (STRMK, OMIM #185070) [15–18]. Stormorken syndrome, caused by the STIM1 c.910C > T; p.R304W mutation, is a rare multi-organ disease, presenting with tubular aggregate myopathy, fatigue, anemia, thrombocytopenia, thrombocytopenia, miosis, asplenia, ichthyosis, and headache [10,13,14,18–20]. Interestingly, the effect of GoF STIM1 mutations in the skeletal or dental tissues has not been explored in patients with TAM or Stormorken syndrome [13,14,18,20–23].

STIM1\(^{R304W}\) and ORAI1\(^{R304W}\) mouse models have provided insight into the effects of absent SOCE and the resulting cellular Ca\(^{2+}\) dysregulation [24–28]. These mice present with neonatal lethality, where the surviving mice from both knock-out models displayed significantly reduced growth [24,26–28]. In addition, the ORAI1\(^{R304W}\) mice presented with a severe osteoprotic bone phenotype, characterized by a decreased bone volume versus tissue volume and decreased trabecular number and increased trabecular spacing [26].

On the other hand, mouse models expressing STIM1 GoF mutations causing constitutive SOCE also manifested pathological features [29–31]. Heterozygous Stimsax mice expressing the STIM1 EF-hand mutation D84G [30], which causes TAM1 in humans [10], developed spleen abnormalities including increased spleen size and abnormal red and white pulp architecture. The Stimsax mice also show bone marrow fibrosis and increased collagen deposition which affected the femoral bone cavity in heterozygous state [30]. Two mouse models expressing STIM1 R304W have been reported [29,31]. Both models are described with reduced body size, hematological and muscular defects, and homozygous lethality. The STIM1\(^{R304W}\) model described by Silva-Rojas et al. also displayed spleen enlargement, reduced bone marrow cavity, abnormal cortical and trabecular architecture and diminished bone strength [31].

In this paper, we extend the phenotyping of our mouse line expressing the STIM1 R304W mutation, named Stim1\(^{R304W}\) [29], to further elucidate the abnormalities in the skeletal system due to constitutive SOCE. We describe macroscopic skeletal anomalies and severe changes in the bone architecture in the Stim1\(^{R304W}\) mice, as well as a novel feature of ectopic subgingival hair growth on the labial side of the lower incisors.

### 2. Methods

#### 2.1. Maintenance of mice

All animals were housed at standard conditions (12:12-hr light/dark, 21 ± 2 °C temperature and 55 ± 5 % relative humidity) with ad libitum access to water and standard chow. All experimental protocols for generation, breeding and use of transgenic animals in Norway were registered and approved by the Norwegian Food Safety Authority (Mattilsynet, Experimental animal welfare supervision and application system ID 7216 and 6991) and followed the ethical guidelines given in Directive 2010/63/EU of the European Parliament on the protection of animals used for scientific purposes.

The Stim1\(^{R304W}\) mouse model carrying the Stimi\(^{R304W}\) (NC_000073.6) chr7:102424171A > T (GRCM38.p4) mutation, resulting in a R304W amino acid substitution in STIM1 was established by Gamage et al. [29]. The mouse line was generated on a C57bl6xCBA F1 background using Zinc Finger technology [29]. All animals in the study were euthanized at 5–6 months of age and a full necropsy was performed. Tissues were stored in 10 % formalin or snap frozen until further analysis.

#### 2.2. Spleen measurement

Total body weight was measured in live animals before euthanasia. Upon euthanasia, the spleen of each animal was dissected, photographed, weighed and length measured. The size of spleen is presented as a ratio of spleen and total body mass at euthanasia.

#### 2.3. Whole mouse radiographs

Whole animal radiography of formalin fixed mice was performed using Philips X-ray generator and tube (Philips Norge AS, Health Systems, Oslo, Norway) and Sound Eklin DR developer (Medivet Scandinavian AB, Angelholm, Sweden).

#### 2.4. Microcomputed tomography (μCT) analysis of the tibia

Microcomputed tomography (μCT) scanning was performed to measure morphological indices of metaphysial regions of tibias. Formalin fixed bone samples wrapped in plastic foil were rotated around their long axes with a rotation step of 0.4° and images were acquired using Bruker Skyscan 1172 (Bruker, Kontich, Belgium) with the following parameters: pixel size =5 μm; peak tube potential =100 kV; X-ray intensity =100 μA; 0.5-mm aluminum filter. Raw images were reconstructed by the NRecon Reconstruction Software (Microphotonics, Inc. Allentown, PA) to 3-dimensional cross-sectional image data sets, using a 3-dimensional cone beam algorithm. For the reconstruction, beam hardening was set to 0 %, smoothing to 1, and ring artefact reduction to 10. Structural indices were calculated on reconstructed images using the CT-Analyser (CTAn) software package (Blue Scientific Limited, Cambridge, UK). Cortical and trabecular bone were separated using a custom processing algorithm in CTAn, based on the different thicknesses of the structures. Cortical bone was analyzed by a region of 180 slices, starting 9 mm distal to the metaphysis. Cortical parameters included mean total cross sectional area, cortical bone volume and cortical bone area. Trabecular bone was analyzed in the proximal metaphysis region, starting just distal to the metaphysis and continued distally for 240 slices. Trabecular parameters included trabecular separation, trabecular bone volume fraction and trabecular number.

#### 2.5. Histology

Formalin fixed tibiae from Stim1\(^{−/−}\) and Stim1\(^{R304W/R304W}\) mice, were decalcified in a decalcification solution (34 g Sodium formate (HCOONa), 151 ml concentrated formic acid, 11. H\(_2\)O) for 24 h and paraffin embedded. The tissue blocks were then sectioned at 4 μm thickness and stained with hematoxylin and eosin (HE) using standard protocols.

#### 2.6. Immunohistochemistry

Adjacent sections obtained from tissue block sectioning were stained for STIM1 using the avidin-biotin method with the Vectastain...
**Fig. 1.** Thoracic kyphosis and 12 pairs of ribs in the Stim1R304W/R304W mice.
(A–B) Lateral view of a representative Stim1R304W/R304W mouse showing thoracic kyphosis (arrowhead) (A), confirmed by whole body X-ray (B).
(C) Representative image of a chest X-ray from Stim1R304W/R304W mouse showing 12 pairs of ribs.

Elite ABC System (PK-6100, Vector Laboratories, Burlingame, CA, USA). Sections were de-paraffinized and antigen retrieval performed at 92 °C for 5 min, in citrate buffer (0.01 M, pH 6.0) followed by a 5 min incubation in the heated solution. Sections were rinsed in distilled H2O and the process was repeated, followed by 15 min incubation in the heated solution and 15 min incubation in 3 % H2O2 in methanol/distilled H2O, and stored overnight at 4 °C in phosphate buffered saline (PBS). The sections were blocked for 30 min in 1:100 normal goat serum in PBS and stained with 1:400 anti-human STIM1 (LS-C209377, LifeSpan BioSciences, Inc., Seattle WA, USA) for 1 h at room temperature on a rotating table followed by a 30 min incubation with 1:50 biotinylated goat anti-rabbit antibody (Vectastain kit). Samples were washed and treated with avidin (1:50 in PBS) and biotinylated horse-radish peroxidase (1:50 in PBS), for 30 min at room temperature followed by rinse with PBS with 0.05 % Tween 20. The STIM1 signal was developed for 8 min with ImmPACT AEC (SK-4205, Vector Laboratories). The sections were subsequently contrast stained with hematoxylin and mounted in xylene.

2.7. **Macroscopy and scanning electron microscopy (SEM) of mouse dental structures**

All animals aged 5 weeks to 6 months were examined macroscopically for presence of ectopic subgingival hair growth below the lower central incisors. Jaws were dissected at euthanasia at 5–6 months age, stored in formalin and or PBS and shipped at room temperature to NYU College of Dentistry for Backscattered Scanning Electron Microscopy (BSE-SEM) analysis as reported previously [32]. Briefly, mandibles of WT and Stim1R304W mice were extracted and cleared of soft tissues, dehydrated, embedded in PMMA resin before sectioning about 2 mm from the incisor tip. After polishing, samples were imaged in parallel in the BSE-SEM without conductive coating setting contrast and brightness to the WT sample.

2.8. **Re-analysis of X-rays and computed tomography (CT) scans from patients with Stormorken syndrome**

Available radiographs from patients F1 III-1, F2 II-3, F3 II-1 and F4 III-1 described by Mische et al., [13] were collected and assessed retrospectively. Images collected include: computed tomography (CT) of the neck, chest and abdomen of F1 III-1; CT of the head and radiographs of chest, hands and lower extremities in F2 II-3; radiographs of cervical spine and lower extremities in F3 II-1; and finally, CT of the head, chest and abdomen and radiographs of the spine of F4 III-1 (numbers as previously reported). Chest X-rays from patient F1 III-2 described by Nesin et al., [14] and the patient described by Borsani et al. [33], were also assessed.

2.9. **Statistics**

Statistical analyses of μCT parameters were performed using the SPSS Statistics software (IBM Corp. Released 2017. IBM SPSS Statistics for Windows, Version 25.0. Armonk, NY: IBM Corp.). Exact significance was tested based on Mann-Whitney U test (2-tailed) to compare the differences between groups. A p-value < 0.05 was defined as statistically significant. Student’s t-test (unpaired) was used to evaluate significant differences in the spleen to body mass ratios.

3. **Results**

3.1. **Stim1R304W/R304W mice showed thoracic kyphosis and reduced number of ribs**

A knock-in mouse line expressing the STIM1 R304W mutation was established [29]. This resulted in perinatal lethality in homozygous state. Only three homozygous mice survived into adulthood following 25 heterozygous crosses, which in total resulted in 130 offspring. The surviving mice presented with reduced growth and body size compared to age and sex matched wild type littermates [29].

The three homozygous mice were euthanized at 5–6 months of age and stored in formalin before the examination of the skeletons. Macroscopic examination of the mice revealed thoracic kyphosis, which was further documented by whole body X-ray radiographs (Fig. 1A–B). The radiographs also revealed the presence of 12 pairs of ribs in all three Stim1R304W/R304W mice (Fig. 1C), which is in contrast to the 13 pairs of ribs in wild type mice. No other abnormal features were observed on X-ray. Thoracic kyphosis was not detected by macroscopic examination of 91 heterozygous Stim1R304W/+ mice. Further micro-computed tomography (μCT) scans and data analysis did not detect differences in the femur to tibia length ratios in the homozygous mice compared to wild type (data not shown). Therefore, the X-ray and μCT results indicated that the skeleton of the Stim1R304W/R304W mice were smaller, but proportionate in comparison to the wild type mice.
3.2. Patients with the STIM1 R304W mutation did not present with clear skeletal anomalies

Motivated by the skeletal anomalies in the Stim1<sup>R304W</sup> mouse line, we did a retrospective analysis on chest X-ray and CT scans in six patients with Stormorken syndrome [13,14,33]. We did not detect skeletal abnormalities in ribs and spine in patient F1 III-1 and F3 II-1 reported by Misceo et al. [13], in patient F1 III-2 described by Nesi et al. [14], and in the patient described by Borsani et al. [33]. Patient F2 II-3 in Misceo et al. [13], also had a normal spine and normal number of pairs of ribs, but rib number 5 on the right side was bifid anteriorly, and presented with periarticular soft tissue calcifications. Patient F4 III-1 in Misceo et al. [13] had normal vertebrae and number of ribs, but showed a slight thoracic scoliosis of about 25°. Bone density was considered normal in all patients evaluated by CT and or X-ray radiographs, including normal thickness of the cortical bone. However, it should be noted that these patients maintain a wild type STIM1 allele, in contrast to the mice that are homozygous for the Stormorken mutation, which thus are expected to have a more severe phenotype. The subtle skeletal features in these patients call for examination of more patients with Stormorken syndrome or other GoF STIM1 mutations to assess for the presence of skeletal anomalies.

3.3. Abnormal bone architecture in the Stim1<sup>R304W</sup> mouse line

Macroscopic examinations of cross sections of the femurs revealed normal bone marrow cavity in the wild type mice, but narrow, almost absent bone marrow cavity in the three Stim1<sup>R304W/R304W</sup> mice.

In the Stim1<sup>R304W/R304W</sup> mice μCT analyses of the tibiae documented pathogenic changes in trabecular and cortical bone architecture compared to age and sex matched wild type mice (Fig. 2A–I). The homozygous mice showed significantly decreased trabecular separation (p < 0.05), increased trabecular bone volume fraction (p < 0.05) and trabecular number (p < 0.05) (Fig. 2D–F) compared to wild type mice. They also showed a decrease in mean total cross-sectional cortical bone area (p < 0.05), cortical bone volume (p < 0.05) and cortical bone surface area (p < 0.05) (Fig. 2G–I). The heterozygous mice showed an intermediate phenotype between the wild type and the homozygous mice (Fig. 2D–F). Thus, the Stim1<sup>R304W</sup> mouse line presented with pathological features in the skeletal system, which includes decreased cortical bone volume and an increased trabecular bone volume fraction, with thinner and more compact trabeculae (Fig. 2A–I).

Histological examination of HE stained sections of the proximal segment of the tibial diaphysis of wild type mice showed normal ossification with slender bone trabeculae and an abundant bone marrow. In contrast, the Stim1<sup>R304W/R304W</sup> mice presented with reduced bone marrow cavity, resulting in a more compact morphology (Fig. 3A–D).

3.4. STIM1 R304W was expressed in megakaryocytes, osteoblasts and osteocytes

The expression of the STIM1 protein in bone was assessed by IHC on bone sections from tibiae of homozygous and wild type mice. The data showed that the osteoblasts lining the bone trabeculae and the osteocytes embedded in the bone matrix were strongly positive for STIM1 in both wild type and Stim1<sup>R304W/R304W</sup> mice (Fig. 3E and F).

We previously documented lack of STIM1 protein in platelets from the Stim1<sup>R304W/R304W</sup>→WT chimeras and in embryonic liver megakaryocytes from the Stim1<sup>R304W/R304W</sup> mice [29]. In the only three Stim1<sup>R304W/R304W</sup> mice surviving to adulthood, however, bone marrow megakaryocytes stained positive for STIM1 by IHC (Fig. 4). Thus, the mutant protein was detected in the hematopoietic lineage in all the three Stim1<sup>R304W/R304W</sup> mice surviving to adulthood.

3.5. Stim1<sup>R304W/R304W</sup> mice presented with splenomegaly

Stim1<sup>R304W/R304W</sup> mice presented splenomegaly and showed a mean spleen to body mass ratio of 0.0125 (n = 3) while WT was 0.00275 (n = 18) (p < 0.005) (Fig. 5). Splenomegaly is observed in mice with impaired bone marrow activity [34], and was also detected in the Stim1<sup>R304W</sup> mice described by Silva-Rojas et al. [31].

3.6. The Stim<sup>R304W</sup> mouse line presented with bilateral subgingival hair growth on the lower incisors

All three homozygous Stim1<sup>R304W/R304W</sup> mice presented with hair originating from the lateral side of the lower incisors, which became evident within the first two months of life (Fig. 6A). Similar ectopic subgingival hair growth on the lower incisors was also detected in 18/70 (26 %) of the heterozygous Stim1<sup>R304W/+</sup> mice examined.

The color and structure of the incisors of the Stim1<sup>R304W</sup> mouse line was comparable to the wild type littermates. The incisors did not have a ‘chalky’ appearance, which could have indicated abnormal mineralization. Cross sections of incisor enamel from one wild type and one Stim1<sup>R304W/R304W</sup> mouse were imaged in BSE-SEM to assess broad differences in mineralization of the enamel. Wild type and Stim1<sup>R304W/R304W</sup> mics did not show significant mineralization differences (Fig. 6B–C). Together, these data point to a defect in epithelial cell fate in Stim1<sup>R304W</sup> mouse line, not previously described in mice with Stim1 or Orai1 mutations.

4. Discussion

Store-operated calcium channels (SOCE) serve as a main Ca<sup>2+</sup>-regulating mechanism in both bone and dental tissue growth and development in man and mouse. This is evident in patients with Ca<sup>2+</sup>-release-activated Ca<sup>2+</sup> (CRAC) channelopathies, who present with defects in dental enamel mineralization [8]. A dental phenotype with chalky enamel is also observed in Stim1-deficient mice [35,36]. In the bone, Orai1 regulates osteoblast and osteoclast formation and differentiation [37]. The role of STIM1 in these processes is less understood [38]. However, based on the importance of SOCE, it can be hypothesized that STIM1 also plays a critical role in these processes and any disruption in STIM1 function may result in imbalances of bone homeostasis and in dental enamel formation. The Stim1<sup>R304W</sup> mouse model demonstrates that constitutive SOCE due to the GoF Stim1 mutation results in skeletal defects and abnormal epithelial cell fate, thereby making Stim1<sup>R304W</sup> mice a tool for exploring the effects of constitutive SOCE induced Ca<sup>2+</sup> dysregulation in these tissues.

STIM1 GoF mutations are pathogenic in mice, but the cellular and phenotypic effects seem to be different for each mutation [29–31]. Embryonic lethality at <14.5 dpc caused by severe bleeding is described in mice expressing the EF-hand GoF mutation Stim1 D84G [30], while homozygous Stim1 R304W mice died perinatally with no signs of bleeding [29]. Stim1 is important in muscle at early developmental stages [39,40], and both the heterozygous Stim1<sup>R304W/+</sup> mice and the surviving homozygotes showed severe skeletal muscle degeneration [29], indicating that the cause of lethality may be understood by further studies of the muscle pathology in our Stim1<sup>R304W</sup> mice.

Our Stim1<sup>R304W</sup> mice showed macroscopic skeletal anomalies: proportionally small skeleton, reduced number of ribs and kyphosis in Stim1<sup>R304W/R304W</sup> mice. We further documented severe microscopic anomalies in the bone architecture of these mice, with increased trabecular to cortical bone fraction, and almost absent bone marrow cavity. This is in line with the observations in the mouse model described by Silva-Rojas et al. [31], expressing the same mutation, which presented with decreased cellular density, reduced bone marrow cavity and abnormal mechanical bone properties. In contrast, re-analysis of X-ray and CT scans of six patients with Stormorken syndrome expressing...
Fig. 2. μCT analyses of the tibia showed increased trabecular to cortical bone fraction in Stim1R304W mice. (A–C) 3D projections of μCT scanning of tibia from Stim1R304W and wild type mice as indicated. Homozygous mice presented decreased cortical bone volume (B) and increased trabecular bone volume fraction with thinner and more compact trabeculae (C) compared to the wild types.

(D–I) Homozygous mice (n = 3) presented with significantly decreased trabecular separation (D), increased trabecular bone volume fraction (E) and trabecular number (F) compared to wild type mice (n = 6). Significantly decreased mean total cross sectional area (G), cortical bone volume (H) and cortical bone area (I) were also seen in Stim1R304W mice (n = 3) in comparison to age matched wild type mice (n = 6).

Heterozygous Stim1R304W mice (n = 6) showed an intermediate phenotype between wild type and homozygous littermates in all aspects above (D–F).

In all panels, p-values for significance are indicated as *=p < 0.05.


the STIM1 R304W mutation detected only minor and varied anomalies, which may also be observed in otherwise healthy individuals. Skeletal examination of additional patients with Stormorken syndrome is needed to assess if STIM1 GoF mutations cause skeletal anomalies in humans.

We previously showed that STIM1 expression was reduced or undetectable in megakaryocytes and platelets of our heterozygous and homozygous Stim1R304W mice at embryonic stages and in fetal liver chimeras [29]. However, the three homozygous Stim1R304W mice surviving to adulthood showed STIM1 expression in bone marrow megakaryocytes (Fig. 4), documenting expression of the mutant protein in the hematopoietic lineage. Osteoclasts also originate from hematopoietic stem cells, which differentiate to form monocyte precursors that subsequently fuse into mature multinucleated osteoclasts, which mediate bone resorption by decalcifying and degrading the bone matrix [41]. ORAI1 and STIM1 are required for the mononuclear osteoclast fusion, a vital step of the late osteoclast differentiation [42], and Orai1 KO mice show reduced trabecular thickness and reduced cortical ossification [26]. The Stim1R304W mice, however, showed increased trabecular bone volume and number. Because the hematopoietic cells of adult homozygous Stim1R304W mice showed STIM1 expression and because the bone phenotype of homozygous and heterozygous Stim1R304W mice differed from the phenotype of mice lacking ORAI1, it is reasonable to postulate that the mutant STIM1 is also expressed in the osteoclasts in the adult homozygous Stim1R304W mice.

Osteoblasts, which secrete matrix proteins necessary for bone mineralization, originate from mesenchymal stem cells in the bone marrow [43,44]. STIM1 is highly expressed in pre-osteoblasts [44], where Ca²⁺ signaling is essential for their differentiation into mature osteoblasts [45]. We documented expression of the mutant STIM1 R304W protein in the osteoblasts in the homozygous Stim1R304W mice. However, more studies are required into the development and function of both osteoblasts and osteoclasts associated with CRAC channel function to understand the exact mechanism and cell type affected in our Stim1R304W mice leading to the pathogenic phenotype.

Mice naturally have an insufficient total bone marrow volume, and
Fig. 3. HE and IHC staining of sections of proximal tibia from Stim1<sup>R304W/R304W</sup> and wild type mice.

(A–B) Ossification of the proximal tibia of age matched wild type (WT) and Stim1<sup>R304W/R304W</sup> (Hom) mice.

Wild type mice showed normal ossification with slender bone trabeculae and abundant bone marrow (A), while Stim1<sup>R304W/R304W</sup> consistently show increased bone mass with broad bone trabeculae and relatively sparse bone marrow by HE staining (B) (magnification 25×).

(C–D) HE stained sections of the proximal tibia from a wild type and Stim1<sup>R304W/R304W</sup>, respectively (magnification 400×).

(E–F) IHC of sections (STIM1) from wild type (E) and Stim1<sup>R304W/R304W</sup> mice (F) showed strongly STIM1 positive osteoblasts (arrows) and embedded osteocytes (arrowheads) (magnification 400×).

**Fig. 4.** STIM1 staining in megakaryocytes from bone marrow of Stim1<sup>R304W/R304W</sup> mice.

IHC of bone marrow showed STIM1 expression in megakaryocytes in adult wild type, Stim1<sup>R304W/+</sup> and Stim1<sup>R304W/R304W</sup> mice (n = 2) (arrowheads) (Magnification: 400×).

Het: heterozygous; Hom: homozygous; WT: wild type.
Ca\textsuperscript{2+} content \cite{35,36}. However, the effect of increased SOCE activity due to overactive STIM1 in dental enamel in mice is not described. Expression of the GoF STIM1 R304W mutation in our Stim\textsubscript{R304W} mice did not seem to cause obvious mineralization differences when comparing to wild type animals by BSE-SEM of incisor enamel. This is in line with the previously reported observation of normal enamel mineralization in mice with increased SOCE due to knock-down of \textit{Orai2} \cite{25}. Together these observations indicate that lack of SOCE has more substantial pathogenic impact on the enamel mineralization than constitutive SOCE in mice.

The ectopic subgingival hair structures in the Stim\textsubscript{1R304W} mice do not seem to replace enamel growth, but are rather generated in addition to the normal enamel. Rodent incisors have a stem cell compartment at the base (the cervical loop) enabling continuous growth of the tooth, as cells differentiate into different cell types including ameloblasts \cite{46}. Tooth morphogenesis is similar to that of hair and other ectodermal tissues \cite{46}, but hair growth in gingiva is uncommon. However, a similar ectopic hair growth as described in our Stim\textsubscript{1R304W} mice was reported in the epithelium specific mediator complex subunit 1 (\textit{Med1}) knock-out mouse model \cite{47}. Yoshizaki et al. \cite{47} suggested that in these mice the lack of \textit{Med1} caused reduced Notch signaling; a pathway which regulates cell fate in different tissues including the dental epithelial cells \cite{48}. With reduced Notch signaling, the dental epithelium converted into epidermal epithelia due to failure in committing to a dental lineage. This in turn caused defects in the development of enamel, while promoting hair growth in \textit{Med1} KO mice \cite{47,48}. Furthermore, epidermal fate is induced by Ca\textsuperscript{2+} \cite{47}. However, elevated Ca\textsuperscript{2+} is also known to downregulate Notch signaling causing defects in cell lineage commitment \cite{49}. Thus, dysregulation of the Ca\textsuperscript{2+} driven epithelial cell fate may be the cause of the ectopic hair growth in the Stim\textsubscript{1R304W} mice. The ectopic subgingival hair development suggest a potential suppression of the Notch pathway in Stim\textsubscript{1R304W/R304W} mice. It is tempting to hypothesize that a similar suppression of the Notch pathway in mesenchymal progenitor cells, osteoblasts, osteocytes and/or osteoclasts, which express STIM1 and are all known to be affected by the Notch signaling \cite{50}, could explain part of the skeletal phenotype of these mice.

5. Conclusion

The Stim\textsubscript{1R304W} mouse line presented a skeletal phenotype characterized by significant pathogenicity in trabecular and cortical bone, kyphosis, reduced number of ribs, and reduced bone marrow cavity. In addition, the mice showed splenomegaly and ectopic growth of subgingival hair possibly caused by a defect in epithelial cell fate. Our data further confirm the previous reports that elevated SOCE does not seem to affect enamel mineralization \cite{25}, but cause severe defects in bone architecture in mice \cite{30,31}. This further highlights the importance of the ER-protein STIM1 in skeletal system development and homeostasis. Further studies of the osteoclasts, osteoblasts, ameloblasts and epithelial stem cells expressing the mutant STIM1 R304W protein may reveal the dysregulated molecular pathways that cause the observed abnormalities in the Stim\textsubscript{1R304W} mouse line.

Declaration of Competing Interest

There are no conflicts of interests to declare.

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Fig. 6. Ectopic subgingival hair growth in the Stim1R304W mouse line.

(A) Representative image from a Stim1R304W/R304W mouse shows bilateral ectopic hair growth from the labial side of the lower incisors appearing to originate from the gingiva. Macroscopic enamel morphology appeared normal, without chalky appearance.

(B–C) BSE-SEM of cross section of incisors did not reveal differences in signals (brightness) between the wild type (B) and the Stim1R304W/R304W mouse (C).

References


