Loss of DDRGK1 modulates SOX9 ubiquitination in spondyloepimetaphyseal dysplasia

Adetutu T. Egunsola,1 Yangjin Bae,1 Ming-Ming Jiang,1 David S. Liu,1 Yuqing Chen-Evenson,1 Terry Bertin,1 Shan Chen,1 James T. Lu,2,3 Lisette Nevarez,4 Nurit Magal,5 Annick Raas-Rothschild,6,2 Eric C. Swindell,8 Daniel H. Cohn,4,9,10 Richard A. Gibbs,1,2 Philippe M. Campeau,11 Mordechai Shohat,12 and Brendan H. Lee1

1Department of Molecular and Human Genetics, 2Human Genome Sequencing, and 3Department of Structural and Computational Biology and Molecular Biophysics, Baylor College of Medicine, Houston, Texas, USA. 4Department of Molecular Cell and Developmental Biology, UCLA, Los Angeles, California, USA. 5Recanati Institute of Medical Genetics at Rabin Medical Center, Petach Tikva, Israel. 6Institute for Rare Diseases at Sheba-Tel Hashomer Medical Center, Ramat Gan, Israel. 7Department of Pediatrics, Sackler School of Medicine, Tel Aviv University, Tel Aviv, Israel. 8The University of Texas Graduate School of Biomedical Sciences, Houston, Texas, USA. 9Department of Orthopaedic Surgery and 10International Skeletal Dysplasia Registry, UCLA, Los Angeles, California, USA. 11Department of Pediatrics, University of Montreal, Montreal, Quebec, Canada. 12Maccabi Genetic Institute and Bioinformatics Unit — Sheba Cancer Research Center, Sackler School of Medicine, Tel Aviv University, Tel Aviv, Israel.

Shohat-type spondyloepimetaphyseal dysplasia (SEMD) is a skeletal dysplasia that affects cartilage development. Similar skeletal disorders, such as spondyloepiphyseal dysplasias, are linked to mutations in type II collagen (COL2A1), but the causative gene in SEMD is not known. Here, we have performed whole-exome sequencing to identify a recurrent homozygous c.408+1G>A donor splice site loss-of-function mutation in DDRGK1 domain containing 1 (DDRGK1) in 4 families affected by SEMD. In zebrafish, ddrgk1 deficiency disrupted craniofacial cartilage development and led to decreased levels of the chondrogenic master transcription factor sox9 and its downstream target, col2a1. Overexpression of sox9 rescued the zebrafish chondrogenic and craniofacial phenotype generated by ddrgk1 knockout, thus identifying DDRGK1 as a regulator of SOX9. Consistent with these results, Ddrgk1−/− mice displayed delayed limb bud chondrogenic condensation, decreased SOX9 protein expression and Col2a1 transcript levels, and increased apoptosis. Furthermore, we determined that DDRGK1 can directly bind to SOX9 to inhibit its ubiquitination and proteasomal degradation. Taken together, these data indicate that loss of DDRGK1 decreases SOX9 expression and causes a human skeletal dysplasia, identifying a mechanism that regulates chondrogenesis via modulation of SOX9 ubiquitination.

Introduction

Chondrodysplasias are a clinically and genetically diverse group of skeletal dysplasias that primarily affect cartilage and present with disproportionate short stature and premature osteoarthritis (OA) (1–3). Shohat type spondyloepimeta- physeal dysplasia (SEMD) is a chondrodysplasia characterized by vertebral, epiphyseal, and metaphyseal abnormalities. These abnormalities include scoliosis with vertebral compression fractures, flattened vertebral bodies, and hypomineralization of the long bones. Additionally, affected individuals may have a small trunk, short neck, small limbs, joint laxity, bowlegs, and/or abdominal distension with hepatosplenomegaly (4–6). Shohat-type SEMD shares some clinical features with the spondyloepiphyseal dysplasia (SED) spectrum, including long-bone and vertebral defects. Disorders of the SED spectrum, which include SED congenita and X-linked SED tarda, are usually caused by mutations altering the expression and/or processing of type II collagen (COL2A1), the most abundant extracellular matrix protein in cartilage (1, 2, 7, 8). Specifically, mutations in COL2A1 were the first reported cause of a human chondrodysplasia, SED congenita. Shohat-type SEMD is not caused by any known mutations in genes that affect the structure or trafficking of COL2A1. However, its phenotypic overlap with the SED spectrum led us to hypothesize that its causative gene may control COL2A1 processing and/or cartilage formation.

To identify the genetic cause of Shohat-type SEMD, we performed whole-exome sequence (WES) on a proband and identified a homozygous c.408+1G>A donor splice site mutation in the DDRGK domain containing 1 (DDRGK1) gene. DDRGK1, also known as UFM1-binding protein 1 containing a PCI domain (UFBP1), has been linked to protein stability and activation. For instance, in vitro knockdown of Ddrgk1 indirectly inhibits IκBα ubiquitination by increasing phosphorylation of IκBα, a prerequisite for its ubiquitination (9). DDRGK1 is also required for estrogen receptor α (ERα) transactivation, as it mediates activating signal cointegrator 1 (ASC1) ufmylation, an ubiquitin-like posttranslational modification (10). While the cellular functions of Ddrgk1 have been investigated in other systems (9–13), its role in skeletal development remains unknown.

Here, we report that Ddrgk1 is essential for cartilage development and the differentiation of chondrocytes, the cells that produce and maintain cartilage. Furthermore, we demonstrate that DDRGK1 loss of function causes Shohat-type SEMD pathogenesis by increasing ubiquitin-dependent proteasomal degradation of SOX9.

Results

Shohat-type SEMD is caused by a loss-of-function mutation within the DDRGK1 gene. Family 1, of Jewish Iraqi descent, was first
The patients in family 1 (Figure 1, C and D). In addition to severe scoliosis, vertebral compression fractures, platyspondyly, broadened hypomineralized metaphyses, and hypomineralized epiphyses (Figure 1E), patient 5 had a narrow trachea and severe upper airway obstruction requiring a tracheostomy (Figure 1E) with continuous positive airway pressure (CPAP) therapy while sleeping.

To identify the genetic cause of Shohat-type SEMD, we performed WES on 3 affected individuals from the first 2 Jewish families of Iraqi descent (Figure 1, A and C). By applying WES with an established method and variant filtering (14), we identified 1,951 genes with rare coding variants, 53 of which were shared by all 3 individuals. Out of the 53 genes, 13 had homozygous variants in all 3 patients; among these, \textit{SLC4A11} and \textit{DDRGK1} had rare variants having minor allele frequencies of less than 0.05 and were located at the same region of chromosome 20. The \textit{SLC4A11} variant found in our patients is a known variant corresponding to dbSNP rs141705330, for which some individuals in the general population in the Exome Aggregation Consortium (ExAC; http://exac.

reported and diagnosed in 1993 with Shohat-type SEMD (Figure 1A). As previously described, these patients have a disproportionate short stature, short neck, small chest, mild abdominal distension with hepatosplenomegaly, lordosis, short limbs, genu varum, and joint hypermobility (6). Radiographically, they have delayed epiphyseal ossification, platyspondyly with central notches in the vertebral end plates, radiolucency of the femoral metaphyses, and relative fibular overgrowth (6), as seen in the radiographs of patients 1 and 2 at ages 9 months and 5 years, respectively (Figure 1B). At age 18, because of his narrowed trachea, patient 1 required a lifesaving tracheostomy during an aborted orthopedic surgery. Unfortunately, this surgical complication was the cause of death for patients 3 and 4 from families 1 and 4, respectively. In addition, now at age 26, patient 1 has joint pain and hoarseness due to his severely narrowed larynx and bronchi.

The affected individuals in families 2 through 4, of Jewish Iraqi descent, were diagnosed with very short limbs in utero, and at birth, had vertebral and long-bone defects similar to those of

Figure 1. Homozygous \textit{DDRGK1} loss-of-function mutation identified in Iraqi Jewish families with Shohat-type SEMD. (A) Pedigree of family 1. M represents mutant allele. Asterisks indicate individuals who were selected for WES. (B) Radiographs from family 1 reveal platyspondyly and hypomineralized epiphyses and metaphyses in patients 1 and 2, respectively. (C and D) Pedigrees of families 2, 3, and 4. (E) Radiographs show severe scoliosis, vertebral compression fractures, platyspondyly, broadened hypomineralized metaphyses, and smaller than average hypomineralized epiphyses in patient 5 from family 4. (F) Schematic of the c.408+1G>A mutation in the \textit{DDRGK1} gene. (G) Western analysis of whole cell lysates reveals absence of \textit{DDRGK1} expression in patient LCLs.

broadinstitute.org/database) were homozygous. SLC4A11 defects cause corneal endothelial dystrophy and sensorineural hearing loss in humans and mice (15, 16), while DDRGK1 has been linked to protein stability and activation (9, 10). Therefore, based on the known variant and function of SLC4A11, DDRGK1 appeared to be the best candidate gene for Shohat-type SEMD.

The 3 affected individuals had a homozygous c.408+1G>A donor splice site mutation in the third intron of the DDRGK1 gene, while none of the unaffected family members were homozygous for this mutation (Figure 1F). Additionally, by conventional Sanger sequencing, we identified 2 more patients from families 3 and 4 who were homozygous for the same mutation (Figure 1, C and D). This variant was not found in the ExAC database, but there were other potential DDRGK1 deleterious variants in the ExAC database, all heterozygous with minor allele frequencies of less than 0.001.

To assess whether the c.408+1G>A mutation affected DDRGK1 transcription and translation, we performed quantitative real-time PCR (qRT-PCR) and Western blot analysis on RNA and whole-cell lysates from lymphoblastoid cells (LCLs) of patient 1. While control LCLs expressed WT DDRGK1 RNA, patient LCLs expressed a mix of 2 aberrant DDRGK1 RNA species (Supplemental Figure 1, A and B; supplemental material available online with this article; https://doi.org/10.1172/JCI90193DS1). The first aberrant RNA was a read-through of the third intron and the second one was the result of use of a cryptic splice site, both resulting in a premature stop codon (Supplemental Figure 1B). When we examined protein expression, LCLs from the patient did not express DDRGK1 (Figure 1G). These results suggest that the c.408+1G>A homozygous donor splice site mutation in DDRGK1 leads to nonsense-mediated decay and is a founder, disease-causing variant of Shohat-type SEMD.

Knockdown of ddrk1 disrupts cartilage development in zebrafish embryos. In zebrafish, ddrk1 is highly expressed in the head at 60 hours post fertilization (h.p.f.) and the sequence of its Ddrgk1 protein is 53% identical to its human protein ortholog (17). Therefore, zebrafish were used to evaluate the functional consequences of ddrk1 deficiency in cartilage development. During zebrafish embryogenesis, chondrogenesis starts as early as 52 h.p.f., and well-developed craniofacial cartilage structures can be visualized by Alcian blue staining by 120 h.p.f. (18–20). We transiently knocked down the ddrk1 in zebrafish embryos with morpholino oligonucleotide (MO) injections. Similarly to WT, approximately 97% of embryos injected with control MO (control morphants) had well-developed Meckel’s cartilage, ceratohyal cartilage, ceratobranchial cartilage, and neurocranial cartilage by 120 h.p.f. (Figure 2, A and B). Interestingly, 50% to 61% of embryos injected with ddrk1 MO showed craniofacial defects, which increased in severity in a ddrk1 knockdown manner (Figure 2, A and B). These defects included shortened and poorly developed Meckel’s cartilage and ceratobranchial cartilage. Embryos lacking ceratobranchial cartilage showed mild craniofacial defects, while embryos with several craniofacial defects had no neurocranial or ceratobranchial cartilage and poorly developed Meckel’s cartilage and ceratohyal cartilage. WT, n = 44; 5 pg control MO, n = 36; 3 pg ddrk1 MO, n = 50; 4 pg ddrk1 MO, n = 47; 5 pg ddrk1 MO, n = 43; 5 pg ddrk1 MO + 25 pg ddrk1 mRNA, n = 45; 5 pg ddrk1 MO + 50 pg ddrk1 mRNA, n = 38. *P < 0.05; ***P < 0.001; Kruskal-Wallis rank-sum test followed by Wilcoxon’s rank-sum test with continuity correction.
with normal craniofacial structures increased from 51% to 68% (Figure 2, A and B). This rescue of the cartilage phenotype in ddrgk1 morphants by overexpression of ddrgk1 mRNA confirms that ddrgk1 is essential for cartilage development in zebrafish and supports the hypothesis that the c.408+1G>A DDRGK1 loss-of-function mutation causes Shohat-type SEMD.

Ddrgk1−/− mice show delayed chondrogenic mesenchymal condensation in the limb buds. To further characterize the in vivo function of Ddrgk1 in a mammalian context, we generated Ddrgk1 knockout mice using CRISPR/Cas9. After injecting Cas9 mRNA and Ddrgk1 guide RNA into mouse embryos, we identified a 310-bp c.-249_61del mutation in the Ddrgk1 gene in a founder mouse. This mutation deletes a fragment of the 5' UTR and most of the first exon, including the start codon (Supplemental Figure 2, A and B). RT-PCR and Western blot analysis confirmed that Ddrgk1−/− (Ddrgk1−/−) mice do not express Ddrgk1 mRNA or protein (Supplemental Figure 2, C and D). As previously reported, we confirmed that Ddrgk1 loss of function results in embryonic lethality between E11.5 and E12.5 (ref. 13, Supplemental Figure 2E and Supplemental Table 1) because of an erythropoiesis deficiency (13). Deletion of Ddrgk1 decreases the number of erythrocytes and their progenitor cells in peripheral blood, while increasing the number of abnormal multinucleated erythrocytes in E11.5 embryos (13). However, there are no reports of Shohat-type SEMD patients with any erythropoiesis deficiency or phenotype (5, 6). These results suggest that erythropoiesis deficiency is potentially a mouse-specific phe-
notype, which would explain why embryonic lethality occurs in Ddrgk1<−− mice and not in Shohat-type SEMD patients.

To evaluate the consequences of Ddrgk1 loss during chondrogenesis, we assessed chondrogenic mesenchymal condensation from Ddrgk1−/− limb buds at E11.5 and E12.5. We observed proper patterning of digit formation in WT limb buds at E11.5 and E12.5 by H&E staining (Figure 3, A and B). However, condensation of mesenchymal cells did not occur in Ddrgk1−/− limb buds up to E12.5, which suggests that Ddrgk1 deletion causes a delay or absence of the early stages of chondrogenesis (Figure 3, A and B). These results are consistent with those obtained from ex vivo chondrogenic differentiation assay. After 7 days, E11.5 Ddrgk1−/− mesenchymal cells generated fewer Alcian blue–positive cartilaginous nodules than E11.5 WT cells (Figure 3C). Additionally, deletion of Ddrgk1 increased apoptosis and cell death, trending toward significance, in E11.5 limb buds (Figure 3D). Taken together, the phenotypes of ddrgk1 morphants and Ddrgk1 null mouse indicate that Ddrgk1 is required for chondrocyte differentiation from mesenchymal cells, and furthermore, they suggest that loss of DDRGK1 reduces the chondrogenic potency of mesenchymal cells, causing the cartilage phenotype seen in Shohat-type SEMD.

Figure 4. Ddrgk1 deficiency decreases SOX9 protein and Col2a1 mRNA expression. (A and B) ATDC5 cells were transiently transfected with control or Ddrgk1 siRNA and treated with either DMSO or ITS to induce differentiation 24 hours later. The cells were harvested 7 days after treatment. (A) RT-PCR of total RNA from ATDC5 cells treated with control siRNA + DMSO (n = 3), Ddrgk1 siRNA + DMSO (n = 3), control siRNA + ITS (n = 3), and Ddrgk1 siRNA + ITS (n = 3). Values are represented as mean ± SEM. **P < 0.01, 2-way ANOVA followed by Tukey’s post-hoc test. (B) Immunoblots of total cell lysates from ATDC5 cells. The immunoblots are representative of 3 independent experiments. (C) ddrgk1 morphants have less mRNA expression of col2a1, but not of sox9, than control morphants. Zebrafish embryos were injected with 5 pg control MO (n = 3) or 5 pg ddrk1MO (n = 3), and total RNA was collected for RT-PCR 72 h.p.f. later. Values are represented as mean ± SEM. **P < 0.01, **P < 0.001, 2-tailed t test. (D and E) Deletion of Ddrgk1 decreases transcript levels of Col2a1 via SOX9 protein reduction in E11.5 limb buds. (D) RT-PCR of total RNA from E11.5 WT (n = 3) and Ddrgk1−/− (n = 3) limb buds. Values are represented as mean ± SEM. *P < 0.05, 2-tailed t test. (E) Immunoblots of total cell lysates from E11.5 WT (n = 3) and Ddrgk1−/− (n = 3) limb buds.
and Ddrgk1−/− mice. At 72 h.p.f., ddrk1 morphants had decreased transcript levels of col2a1-1a and col2a1-1b, but not sox9a or sox9b, compared with control morphants (Figure 4C). Also, at E11.5, Ddrgk1−/− mice had decreased protein expression of Sox9 and reduced Col2a1 mRNA levels in their limb buds (Figure 4, D and E). Overall, these results demonstrate that Sox9 is a downstream target of DDRGK1.

To further validate sox9 as a downstream target of ddrk1 in zebrafish, we performed rescue experiments overexpressing sox9a in ddrk1 zebrafish morphants. Overexpressing sox9a mRNA caused 60% lethality in zebrafish embryos by 120 d.p.f. (data not shown), which is consistent with the 80% postnatal lethality in Col2a1+/-/ mice overexpressing Sox9 in the chondrocytes (24). When we coinjected ddrk1 MO with sox9a mRNA into zebrafish embryos, it rescued the ddrk1 craniofacial defects in a dosage-dependent manner (Figure 5, A and B). Only 43% of ddrk1 morphants had normal craniofacial features; however, as we increased sox9a mRNA in ddrk1 morphants from 50 pg to 150 pg, the proportion of ddrk1 morphants with normal craniofacial structures increased from 48% to 75% (Figure 5B). The rescue of the ddrk1 cartilage phenotype by sox9a overexpression confirms sox9 as a downstream target of ddrk1. These results suggest that in Shohat-type SEMD patients, the c.408+1G>A DDRGK1 loss-of-function mutation exerts its effects by decreasing SOX9 in the skeleton.

**Discussion**

In this study, we identified a homozygous c.408+1G>A donor splice site mutation in DDRGK1 as a genetic cause of Shohat-type SEMD. Specifically, this mutation causes aberrant splicing and a loss of DDRGK1 in patient LCLs. Similarly to what is observed in humans, we demonstrated that Ddrgk1 deficiency disrupts chondrogenesis and cartilage formation in mice and zebrafish; however, unlike in Shohat-type SEMD patients, loss of Ddrgk1 in mice is embryonic lethal. Though previous studies have shown that Ddrgk1 deficiency causes defective erythropoiesis in mice (13),
we and others have not observed this phenotype in patients with Shohat-type SEMD. At the same time, a GWAS identified a link because of its essential role in regulating multiple developmental pathways, including chondrogenesis, gonadogenesis, gliogenesis, redundant pathways and to assess whether they also have a role in human phenotypes, perhaps due to physiological functional redundancy in mice. In the future, it would be interesting to identify these redundant pathways and to assess whether they also have a role in chondrogenesis during cartilage development.

During development, there are numerous master regulators that orchestrate and control central tissue-specific developmental pathways. To list a few, RUNX2, MYOD, and TAL1 are master transcription factors that regulate the differentiation of osteoblast, skeletal muscle, and erythrocytes, respectively (28–30). On the other hand, SOX9 is a unique master transcription factor that orchestrates and controls central tissue-specific development. Investigating Shohat-type SEMD thus identified changes in expression of SOX9 and its downstream target gene, Col2a1. Using CRISPR/CAS9-generated Ddrgk1−/− mice and differentiated ATDC5 cells, we found that Ddrgk1 deficiency decreases Sox9 protein levels and the expression of its target gene, Col2a1. We then rescued the craniofacial phenotype in ddrk1 knockdown zebrafish by overexpressing sox9a to confirm that Sox9 is an important epistatic downstream target of Ddrgk1. Though the reduction in Sox9 protein levels is less than 50% in Ddrgk1−/− mice, these mice have a more severe skeletal phenotype than Sox9−/− mice haploinsufficient for Sox9 (38). One possible explanation for the more severe phenotype in Ddrgk1−/− mice may be that DDRGK1 regulates multiple targets involved in chondrogenesis in addition to SOX9. Another interesting finding that requires further investigation is that knockdown of Ddrgk1 decreases SOX9 in differentiated ATDC5 cells, but not in undifferentiated ATDC5 cells. These results suggest that the control of SOX9 protein stability by DDRGK1 is temporally regulated. The function of this potential temporal regulation, the factors that govern this regulation, and how they are regulated are interesting subjects for future studies.

The clinical overlap among CMD, the SED spectrum of disorders, and Shohat-type SEMD supports that their pathogenesis is caused by mutations in the same pathway. The SED spectrum phenotypes are caused by mutations altering the structure and trafficking of COL2A1, whereas CMD and Shohat-type SEMD are both caused by mutations affecting SOX9 in its regulation of COL2A1. Our study emphasizes the importance of studying phenocopies of disease because of the potential to reveal novel components of a developmental pathway. Investigating Shohat-type SEMD thus identified DDRGK1 as an indirect regulator of COL2A1 expression through its ability to inhibit SOX9 ubiquitin-dependent proteasomal degradation.

In the present study, we show that DDRGK1 interacts in a complex with SOX9 to regulate its ubiquitination; however, the underlying mechanism mediating this inhibition remains unknown. DDRGK1 is known to be required for umfylation, a ubiquitin-like posttranslational modification that covalently binds ubiquitin-like fold modifier 1 (UFM1) to substrates (10, 12, 40, 41). DDRGK1 forms a complex with UFM1 substrates and UFM1-specific ligase 1 (UFL1), an UFM1 E3 ligase, and DDRGK1 knockdown prevents ASC1 umfylation (10, 12). Therefore, based on these studies and our findings, it is possible that Sox9 may be umfylated, a process for which DDRGK1 is required; SOX9 umfylation in turn may prevent its ubiquitination and proteasomal degradation. Alternatively, another potential mechanism for DDRGK1 inhi-
bition of SOX9 ubiquitination may be independent of its role in ubiquitylation. DDRGK1 has a proteasome, COP9, initiation factor 3 (PC1) domain (ref. 42 and Figure 1F), which can also be found in large multisubunit protein complexes, including the proteasome, COP9 signalosome, and eukaryotic translation initiation factor 3 (43–45). The PCI domain facilitates protein-protein interactions, and it is thought to facilitate interactions in complexes containing this domain (43–45). Therefore, DDRGK1 may interact with the proteasome complex to directly mediate SOX9 ubiquitin-dependent proteasomal degradation.

In conclusion, our results demonstrate that the homozygous c.408+1G>A loss-of-function mutation in DDRGK1 causes Shohat-type SEMD by loss of DDRGK1 inhibition of SOX9 ubiquitin-dependent proteasomal degradation, thus impairing chondrogenesis. By using a rapid multimodel system (a combination of morpholino-knockdown zebrafish, CRISPR/Cas9-generated mouse genetic models, and in vitro assays) to study the pathogenicity of a rare genetic variant, we identified the genetic cause of Shohat-type SEMD. Previous studies of DDRGK1 investigated its cellular and in vivo functions in mice, but our study addressed, for what we believe is the first time, its physiological role in the context of a human disease and skeletogenesis. Furthermore, this work reveals a mechanism regulating the developmental pathway of chondrogenesis.

Methods

WES. Total genomic DNA was extracted from whole blood from 3 affected individuals. WES was performed on 3 affected individuals from families 1 and 2 and the data analyzed, as previously described (14). Afterwards, conventional Sanger sequencing was performed on all living patients to confirm the DDRGK1 variant identified in the WES analysis, as previously described (14).

Cell culture, transfection, and treatment. LCLs were extracted from the proband and a control who did not have Sutcliffe type of spondyloepiphyseal dysplasia. LCLs were grown at 37°C in αMEM (HyClone) plus 10% FBS and with 100 units/ml of penicillin and 1 μg/ml of streptomycin. HEK293T cells (ATCC) were grown at 37°C in DMEM (HyClone) supplemented with 100 units/ml of penicillin, 1 μg/ml of streptomycin, and 10% FBS. 293T cells were transiently transfected with expression plasmids for 48 hours, using Lipofectamine 2000 (Invitrogen) according to the manufacturer’s protocol. ATDC5 cells (ATCC) were grown at 37°C in DMEM/F12 (HyClone) supplemented with 100 units/ml of penicillin, 1 μg/ml of streptomycin, and 5% FBS. For the in vitro knockdown experiments, ATDC5 cells were transiently transfected with either 50 nM siGENOME mouse Ddrgk1 siRNA or 50 nM siGENOME Non-Targeting siRNA Pool #1 (Dharmacon) using Lipofectamine 2000 (Invitrogen). To differentiate ATDC5 cells, cells were treated 24 hours after siRNA treatment with 1× ITS Liquid Media Supplement (100×, Sigma-Aldrich) for 7 days. For the ubiquitination assays, 293T cells were treated with 20 μM of proteasome inhibitor MG132 6 hours before harvesting. To inhibit translation, 293T cells were treated with 10 μg/ml of protein synthesis inhibitor cycloheximide for 6 hours before harvesting.

Antibodies. The rabbit anti-DDRGK1 antibody was a gift from Frans Shuit (12). Anti-SOX9 antibody (catalog AB5535), anti–c-Myc antibody (catalog sc-40), anti-FLAG antibody (catalog F1804), anti-BrdU antibody (catalog Ab1893), and normal rabbit IgG (catalog 2729) were purchased from Millipore, Santa Cruz Biotechnology Inc., Sigma-Aldrich, Abcam, and Cell Signaling Technology, respectively. The anti-SOX9 antibody (AB5535) and anti-BrdU antibody (AB1893) were purchased from Abcam. The secondary antibodies used were goat anti-mouse IgG-HRP (H+L) conjugate (catalog 172011, Bio-Rad), goat anti-mouse antibody (catalog 926-68070, Licor), goat anti-rabbit antibody (catalog 926-32211, LiCor), donkey anti-sheep antibody Alexa Fluor 594 (catalog Abi50180, Abcam), and anti-rabbit IgG, HRP-linked whole antibody (catalog NA934, GE Healthcare Life Sciences).

Expression plasmids. We previously generated the pcDNA3.1.FLAG-SOX9 plasmid (46); the pcMV_Myc-DDRGK1 and His6UB plasmids were gifts from Honglin Li (11) and Dae-Sik Lim (47), respectively. The ddrkgk1 and sox9a CDNs were amplified from zebrasfish CDN using the primers listed in Supplemental Table 2 and then cloned into the pCS2 vector by In-Fusion Cloning (Clontech). Afterwards, to prevent ddrkgk1 MO targeting of exogenous ddrkgk1 mRNA, 5 nucleotide sequences of ddrkgk1 pCS2 vector were mutated by QuickChange (Agilent Technologies) site-directed mutagenesis without altering the protein sequence (Supplemental Table 2).

Microinjection and Alcian blue staining of zebrafish embryos. From 3 to 5 pg ddrkgk1 translational MO or ddrkgk1 control MO (Gene Tools) was injected into embryos at the 1-cell stage. For rescue experiments, mutated ddrkgk1 mRNA and sox9a mRNA were transcribed from the pCS2 vectors using the mMessage mMachine SP6 kit (Ambion). Amounts ranging from 25 to 150 pg were used for the injections (Supplemental Table 2). At 120 h.p.f., embryos were fixed in 2% PFA and then stained in 0.1% Alcian blue solution at 4°C for 24 hours. After 24 hours, the embryos were bleached in 3% H2O2 and 0.5% KOH.

CRISPR/Cas9 generation of Ddrgk1−/− mice. Ddrgk1 sgRNA (Supplemental Table 2) was designed using the online CRISPR Design Tool and cloned into the hSpCas9 vector as previously described (48). After cloning, the sgRNA template was generated by PCR amplification, which simultaneously added a T7 promoter in front of the Ddrgk1 sgRNA sequence (Supplemental Table 2). RNA was transcribed from the template using the MEGAscript T7 Transcription Kit (Ambion) and cleaned up with the MEGAClear Transcription Clean-Up Kit (Ambion). 17 ng/μl Ddrgk1 sgRNA and 50 ng/μl Cas9 mRNA were co-injected into C57/B6J mouse embryos (Jackson Laboratory). The founder mice from these injections were screened for mutations in the Ddrgk1 gene (Supplemental Table 2), and the desired line was backcrossed 5 generations.

Histology of mouse limb buds. E11.5 and E12.5 limb buds were collected and fixed in 4% PFA at 4°C for 24 hours. The limb buds were then embedded in paraffin and sectioned 7-μm thick. To examine morphology, sections were stained with H&E (Sigma-Aldrich). To assess apoptosis, sections were stained with the ApopTag Fluorescein In Situ Apoptosis Detection Kit (Millipore) according to the manufacturer’s protocol.

Coimmunoprecipitation. 293T cells were transiently transfected with 3 μg FLAG-Sox9 vector and 3 μg Myc-DDRGK1. After 48 hours, the cells were lysed in lysis buffer (50 mM Tris HCl, pH 8, 150 mM NaCl, 0.1% NP-40, 1 mM EDTA, 2 mM N-ethylmaleimide and proteinase inhibitor cocktail; GenDEPOT), vortexed, and then incubated on ice for 20 minutes. After centrifugation at 13,000 g for 20 minutes at 4°C, the supernatant was preclarified by adding protein G agarose (Roche) and rotating at 4°C for 20 minutes. The supernatant was then incubated with anti-MYC antibody (Sigma-Aldrich) or normal IgG (Santa Cruz Biotechnology Inc.) and bound to protein G beads (Roche). After washing the immunoprecipitated samples 3 times with lysis buffer,
proteins were separated by SDS-PAGE, and SOX9 was detected with anti-FLAG antibody.

**Ubiquitination assay.** 293T cells were transiently transfected with 2 μg FLAG-SOx9 vector, His-UB vector, and 4 μg Myc-DDRGK1 plasmid. After 48 hours, the His-tagged ubiquitinated proteins were pulled down with Ni-NTA agarose beads (QIAGEN) as previously described (49). Following Ni-NTA pulldown, proteins were separated by SDS-PAGE and SOX9 was probed with anti-SOX9 antibody.

**qRT-PCR.** Total RNA was extracted from tissues and cells using TRIzol reagent (Invitrogen) and cDNA was synthesized from total RNA using the SuperScript III First Strand RT-PCR kit (Invitrogen). The qRT-PCR was performed on a LightCycler instrument (Roche).

**Statistics.** All the data are shown as mean ± SEM. Statistical analyses used were a type 2-tailed Student’s t test, 2-way ANOVA analyses used were a type 2-tailed Student’s t test followed by Wilcoxon’s rank-sum test with continuity correction. P < 0.05 was considered significant.

**Study approval.** Prior to their participation in this study, the families provided written, informed consent to participate in the study approved by the IRB at Rabin Medical Center, Petah Tikva, Israel. The families underwent physical examinations, and we obtained radiographs of the patients. WT adult zebrafish strain (AB) and Ddrgk1+/− mice were both maintained under standard laboratory conditions. Zebrafish embryos were obtained after natural crosses between males and females, and mouse embryos were obtained by timed mating. All animal studies were performed under protocols reviewed and approved by IACUCs at the University of Texas Medical School at Houston or Baylor College of Medicine.

**Author contributions**

NM, ARR, and MS were involved in patient care, patient diagnoses, and human data collection. ATE, YB, ECS, PMC, and BHL designed the research studies. JTL, RAG, and PMC conducted the bioinformatics analyses. SC performed the statistical analyses. ATE, YB, MMJ, DSL, YCE, TB, LN, and PMC conducted experiments and acquired data. ATE, YB, ECS, DHC, PMC, and BHL analyzed data. ECS, DHC, and BHL provided the reagents. ATE wrote the manuscript, and YB, TB, ECS, DHC, PMC, MS, and BHL edited the manuscript.

**Acknowledgments**

This work was supported by the Baylor College of Medicine Intellectual and Developmental Disabilities Research Center (HD024064) from the Eunice Kennedy Shriver National Institute of Child Health and Human Development, the Baylor College of Medicine Advanced Technology Cores with funding from the NIH (AI036211, CA125123, and RR024574), the Rolanette and Berdon Lawrence Bone Disease Program of Texas, and the Baylor College of Medicine Center for Skeletal Medicine and Biology. It was also supported by NIH grant R01AR062651 and the Adler chair for Pediatric Cardiology, Tel Aviv University.

Address correspondence to: Yangjin Bae or Brendan H. Lee, Baylor College of Medicine, One Baylor Plaza, R830, Houston, Texas 77030, USA. Phone: 713.798.3548; E-mail: bae@bcm.edu (Y. Bae), blee@bcm.edu (B. H. Lee).


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