# Null and missense mutations of *ERI1* cause a recessive phenotypic dichotomy in humans

# Authors

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Our study uncovers a ribosomopathy-related skeletal dysplasia caused by missense mutations in *ERI1*, which encodes an exoribonuclease. The findings set a precedent showing a more severe effect of missense alleles than null alleles within recessive genetic diseases and imply that ERI1-mediated RNA metabolism regulates development and homeostasis of human cartilage.





# ARTICLE

# Null and missense mutations of *ERI1* cause a recessive phenotypic dichotomy in humans

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### Summary

ERI1 is a 3'-to-5' exoribonuclease involved in RNA metabolic pathways including 5.8S rRNA processing and turnover of histone mRNAs. Its biological and medical significance remain unclear. Here, we uncover a phenotypic dichotomy associated with bi-allelic *ERI1* variants by reporting eight affected individuals from seven unrelated families. A severe spondyloepimetaphyseal dysplasia (SEMD) was identified in five affected individuals with missense variants but not in those with bi-allelic null variants, who showed mild intellectual disability and digital anomalies. The *ERI1* missense variants cause a loss of the exoribonuclease activity, leading to defective trimming of the 5.8S rRNA 3' end and a decreased degradation of replication-dependent histone mRNAs. Affected-individual-derived induced pluripotent stem cells (iPSCs) showed impaired *in vitro* chondrogenesis with downregulation of genes regulating skeletal patterning. Our study establishes an entity previously unreported in OMIM and provides a model showing a more severe effect of missense alleles than null alleles within recessive genotypes, suggesting a key role of ERI1-mediated RNA metabolism in human skeletal patterning and chondrogenesis.

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### Introduction

ERI1 (MIM: 608739) encodes a 3'-to-5' exoribonuclease that consists of an amino-terminal SAP domain (nucleic-acidbinding domain) and a carboxy-terminal 3' exonuclease domain and belongs to the DEDDh family that includes poly(A) ribonuclease, RNase T, and the  $\epsilon$  proofreading subunit of *Escherichia coli* DNA polymerase III.<sup>1</sup> Eri1 is recruited into several conserved and species-specific RNA metabolic pathways. In Caenorhabditis elegans, ERI-1 forms a complex with Dicer that generates specific classes of endogenous siRNAs.<sup>2-4</sup> In Schizosaccharomyces pombe, Eri1 regulates the turnover of chromatin-associated siRNAs.<sup>5</sup> In the mouse, Eri1 targets three types of RNA molecules: rRNA, histone mRNA, and miRNA. Eri1 deficiency in mice results in abnormal 3' end trimming of the 5.8S rRNA, impaired degradation of replication-dependent histone mRNAs, and increased abundance of mature miRNAs.<sup>6–8</sup> The rRNA trimming function of ERI1 is conserved in Caenorhabditis elegans and Schizosaccharomyces pombe.9 However, the impact of ERI1 on these RNA molecules in humans remains unclear.

A homozygous 284 kb genomic deletion, which covers three genes (*ERI1*, *MFHAS1*, and *MIR4660*), was found in an individual with intellectual disability, cardiac malformation, and digit abnormalities.<sup>10</sup> Recently, a homozygous nonsense variant in *ERI1* was reported in an individual with developmental delay and distal limb abnormalities.<sup>11</sup> This evidence supports the involvement of ERI1 in skeletal patterning and implies that a disorder is caused by the disruption of the exoribonuclease function.

In this study, we uncovered a phenotypic dichotomy in eight individuals from seven unrelated families with different types of ERI1 variants and highlighted the association of missense variants with spondyloepimetaphyseal dysplasia (SEMD), which is a group of skeletal diseases characterized by anomalies in spine and long tubular bones. In contrast, the affected individuals with bi-allelic null variants showed mild intellectual disability and digital anomalies. The detailed evaluation of the skeletal phenotypes of Eri1 knockout (KO) mice and the in vitro chondrogenesis of affected-individual-derived induced pluripotent stem cells (iPSCs) supported the functional involvement of ERI1 in skeletal development. Analyses using ERI1 KO HeLa cells and affected-individual-derived cells demonstrated functional conservation of ERI1 with its mouse ortholog in 5.8S rRNA maturation and histone mRNAs decay. The 5.8S rRNA maturation is involved in ribosome biogenesis, defects of which are known to cause ribosomopathies characterized by skeletal dysplasia.<sup>12–15</sup> Our study leads to the findings of an SEMD associated with ribosomopathy and established a framework for understanding the molecular mechanisms underlying the ERI1 phenotypic dichotomy.

### Material and methods

### Next-generation sequencing

The study was approved by the ethical committee of RIKEN (approval number: 17-16-40(6) and data were shared in accor-

dance to those of participating institutions. Genomic DNAs were extracted by standard procedures from peripheral blood of the individuals and their parents after informed consent. Exome sequencing (ES) and genome sequencing were performed as previously described.<sup>16–22</sup> We searched for rare coding or splicing variants in disease-causing genes with transmissions consistent with Mendelian patterns of inheritance.

### PCR and Sanger sequencing

Genomic fragments containing the variants identified by wholeexome sequencing (WES) were amplified by PCR and Sanger sequenced for both strands. The PCR primer sets are shown in Table S1.

### **RNA gel electrophoresis**

Total RNAs were extracted from HeLa cells or the affected-individual-derived lymphoblastoid cells via SV Total RNA Isolation System (Promega). Small RNAs were extracted from HeLa cells via mir-Vana miRNA Isolation Kit (Thermo Fisher Scientific). 2.5  $\mu$ g total RNAs or 3  $\mu$ g small RNAs in Novex TBE-Urea Sample buffer (Thermo Fisher Scientific), Century Plus RNA Marker (Thermo Fisher Scientific), and microRNA Marker (New England Biolabs) were separated on 6% or 15% Novex TBE-Urea Gels in the Novex Mini-Cell (Thermo Fisher Scientific) for 60–70 min at a constant voltage of 180 V. Post-electrophoresis gels were stained by 2  $\mu$ g/mL ethidium bromide for 20 min or SYBR Gold Nucleic Acid Gel Stain (Thermo Fisher Scientific) for 10 min in DEPCtreated water and images were captured under UV light.

#### Construction of expression plasmids

Clones for the coding sequences of *ERI1* (GenBank: NM\_153332.4) were PCR-amplified from a cDNA library derived from human peripheral blood via KOD -Plus- (Toyobo). The PCR amplicons were cloned into the *Sal*1 and *Not*I sites of the pCMV-HA expression vector (Clontech). The mutations were generated by inverse PCR-based site-directed mutagenesis kit (Toyobo). All PCR primers are shown in Table S1. HeLa cells were transfected with TransIT-LT1 (Mirus Bio) according the manufacturer's instructions.

#### Generation of ERI1 KO HeLa cell line

Guide RNA (gRNA) sequences targeting genomic DNA of *ERI1* were designed by E-CRISP (Figure S2A). Complementary oligos containing the gRNA sequences and BbsI ligation adapters were annealed and ligated into the *BbsI*-digested pX330 vector (Addgene #42230).<sup>23</sup> The oligo sequences are listed in Table S1. We co-transfected the pair of pX330 plasmids with pEGFP-N1 (Clontech) vector, which we used as a fluorescent marker to sort transfected cells by FACSAria (BD Biosciences) (Figure S2B). We obtained monoclonal cell populations by limiting dilution of the sorted cells in half 96-well plate as performed previously.<sup>19</sup> The cell lines with deletions covering exon 2 of *ERI1* were identified by PCR for genomic DNA (Figure S2C) and confirmed by Sanger sequencing (Figure S2D). We performed western blot and RNA electrophoresis to validate the knockout effect (Figures S2E and S2F).

#### Western blot

Whole-cell lysates were harvested with 200  $\mu$ L of RIPA Lysis and Extraction Buffer (Thermo Fisher Scientific). Western blots were performed by standard procedures with antibodies for ERI1 (1:100 dilution, sc-137089, Santa Cruz),  $\beta$ -actin-HRP (1:10,000 dilution,

	Individual with	at least one miss	sense variant			Individual wit	th bi-allelic nul	variant		
Individual	Individual 1A	Individual 1B	Individual 2	Individual 3	Individual 4	Individual 5	Individual 6	Individual 7		<i>.</i>
Family	Family 1	Family 1	Family 2	Family 3	Family 4	Family 5	Family 6	Family 7	Hoxha et al. <sup>11</sup>	Choucair et al. <sup>10</sup>
Variant <sup>a</sup>										
DNA	c.[450A>T]; [893A>G]	c.[450A>T]; [893A>G]	c.[464C>T]; [893A>C]	c.[401A>G]; [895T>C]	c.[464C>T]; [62C>A]	c.[514C>T]; [514C>T]	c.[730C>T]; [730C>T]	c.[582+1G>A]; [582+1G>A]	c.[ 352A>T]; [352A>T]	g.[8783887_ 9068578del]; [8783887_ 9068578del]
Protein	p.[Glu150Asp]; [Asp298Gly]	p.[Glu150Asp]; [Asp298Gly]	p.[Pro155Leu]; [Asp298Ala]	p.[Asp134Gly]; [Ser299Pro]	p.[Pro155Leu]; [Ser21*]	p.[Gln172*]; [Gln172*]	p.[Gln244*]; [Gln244*]	p.[?]; [?]	p.[Lys118*]; [Lys118*]	whole gene deletion
Demographics										
Sex	F	М	М	М	М	F	М	F	F	М
Age at last follow-up	28 years	26 years	3.5 months (died)	3.5 months (died)	2 years (died)	8 years 9 months	13 years 3 months	7 years	7 years	5 years 6 months
Weight	24 kg (-5 SD)	26 kg (-5 SD)	NA	3.3 kg (- 4 SD)	failure to thrive	22 kg (8 <sup>th</sup> centile)	62 kg (85 <sup>th</sup> centile)	27.6 kg (50 <sup>th</sup> centile)	normal	N/A
Height	112 cm (-8 SD)	128 cm (-7 SD)	NA	50.3 cm (-5 SD)	short stature	130.8 cm (46 <sup>th</sup> centile)	155 cm (25 <sup>th</sup> centile)	130 cm (90 <sup>th</sup> centile)	normal	105 cm (<3 <sup>rd</sup> centile)
Consanguinity	-	-	-	_	_	-	+	+	_	+
Gestational/perinata	l record									
Fetal ultrasound	hydronephrosis	hydronephrosis	short limbs	N/A	severe IUGR	unremarkable	unremarkable	unremarkable	unremarkable	unremarkable
Gestation age	term	term	term	term	prematurity	term	term	term	term	term
Birth weight	N/A	N/A	2180 g (-3.2 SD)	2,000 g (-3.3 SD)	N/A	2,760 g	2,810 g (-1.6 SD)	2,600 g (-2 SD)	4,010 g (91 <sup>st</sup> centile)	3,000 g (50 <sup>th</sup> centile)
Birth length	N/A	N/A	40 cm (-5 SD)	41.7 cm (-4.3 SD)	N/A	50 cm at 6 weeks (-2.1 SD)	49 cm (-1.3 SD)	normal	49.5 cm (43 <sup>rd</sup> centile)	50 cm (-1.3 SD)
Skeletal radiograph										
Spine anomaly	+	+	+	+	N/A	_	_	-	_	_
Metaphyseal anomaly	N/A	N/A	+	+	N/A	-	_	_	_	_

(Continued on next page)

	Individual with at least one missense variant					Individual with bi-allelic null variant					
Individual Family	Individual 1A Family 1	Individual 1B Family 1	Individual 2 Family 2	Individual 3 Family 3	Individual 4 Family 4	Individual 5 Family 5	Individual 6 Family 6	Individual 7 Family 7	Hoxha et al. <sup>11</sup>	Choucair et al. <sup>10</sup>	
											Epiphyseal anomaly
Syndactyly	+	+	+	+	+	+	-	+	-	+	
Brachydactyly/ clinodactyly/ camptodactyly	+	+	+	+	+	+	+	+	+	+	
Others											
Cardiac anomaly	-	-	+	-	+	+	-	-	-	+	
Hydronephrosis	+	+	N/A	-	+	-	-	+	N/A	-	
Intellectual disability/ developmental delay	_	_	N/A	N/A	$+^{b}$	+ <sup>c</sup>	$+^{d}$	$+^{d}$	+	+	

M, male; F, female; N/A, not available; IUGR, intrauterine growth restriction. <sup>a</sup>Variants are named according to GenBank: NM\_153332.4 or NC\_000008.11. <sup>b</sup>Delayed motor milestones and speech, generalized hypotonia. <sup>c</sup>Global developmental delay, autism. <sup>d</sup>Intellectual disability, mild.



### Figure 1. Photographs of affected individuals

(A and B) Individual 1A. Note zygomatic hypoplasia, prominent alveolar processes of the maxilla and mandible, and small and low-set ears. (C and D) Individual 1B. Note zygomatic hypoplasia and prominent alveolar processes of the maxilla and mandible. (E and F) Maximal elbow extension for individuals 1A and 1B. Note distal camptodactyly in individual 1B.

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PM053-7, MBL, Japan), mouse IgG–peroxidase (1:10,000 dilution, #7076, Cell Signaling Technology), and HA-tag-HRP (1:1,000 dilution, #14031, Cell Signaling Technology). The signals were detected by ECL Prime Western Blotting Detection Reagents (RPN2232, GE Healthcare).

### **Real-time quantitative PCR**

Affected-individual-derived fibroblasts and HeLa cells were treated with 5 mM hydroxyurea for 45 min. Total RNA from affected-individual-derived iPSCs and fibroblasts, and HeLa cells, was extracted with the SV Total RNA Isolation System (Promega) or QIAzol (-QIAGEN) according to the manufacturer's instructions. qPCR was performed as previously described.<sup>19,24</sup> The qPCR primers are listed in Table S1.

### Mice

Mice were housed in specific pathogen-free barrier facilities and used in accordance with protocols approved by the animal care and use committees of the Helmholtz Center, Munich and University of California San Francisco. We created the Eri1-deficient mice by gene targeting in BRUCE4 C57BL/6 ES cells by using standard techniques.7 Wild-type (WT) C57BL/6/J and Actin-FLPe transgenic mice (B6; SJLTg(ACTFLPe)9205Dym/J) were obtained from Jackson Laboratories, CMVCre transgenic mice (6.C-Tg(CMV-cre) 1Cgn) were provided by K. Rajewsky (Harvard Medical School, Boston). To rescue mice from death, which affects offspring on a pure C57BL/6 background,<sup>7</sup> we intercrossed mice once with the NMRI outbred strain background to then generate homozygous F2 mice. The mice used for blood cell tests were on a mixed ICR/ C57BL/6 background, which were similarly generated by crossing ICR mice to  $Eri1^{+/-}$  C57BL/6 mice and backcrossing F1 mice to Eri1<sup>+/-</sup> C57BL/6.<sup>6</sup> Blood was collected in EDTA-coated tubes for red blood cell counts analyzed with a Hemavet system counter (CDC Technologies).

### X-ray examination

Screening of bone phenotypes of the mice was performed at the age of 16 weeks. At the German Mouse Clinic (GMC), mice were maintained in individually ventilated cages with water and standard mouse chow according to German laws and GMC housing conditions.<sup>25</sup> The skeletal morphology of the adult mice was analyzed with Faxitron X-ray Model MX-20 (Specimen Radiography System) equipped with an NTB Digital X-ray Scanner EZ 40 (NTBGmbH). Image processing was performed with ImageJ software.

### Generation of iPSCs

Experimental use of affected-individual-derived cells was approved by ethics committees of RIKEN IMS (Yokohama). Informed consent was obtained from legal guardians of individuals by relevant institutions. The Epstein-Barr virus (EBV)-immortalized lymphoblastoid cell lines (LCLs) were generated from the peripheral blood lymphocytes of the individual and the parents as described previously.<sup>26</sup> LCLs were cultured in Roswell Park Memorial Institute (RPMI) 1640 medium (Sigma) supplemented with 10% fetal bovine serum (Gibco Life Technologies). Reprogram-

ming of LCLs was performed with episomes as previously described with a minor modification.<sup>27</sup> Briefly, LCLs were electroporated with four episomal vectors (pCXLE-hOCT3/4-shp53-F, pCXLE-hSK, pCXLE-hUL, and pCXWB-EBNA1), which were gifts from Shinya Yamanaka (Addgene #27077, #27078, #27080, and #37624).<sup>28</sup> The nucleofected LCLs were seeded on iMatrix-511 (Nippi)-coated plates containing TeSR-E7 reprogramming media (STEMCELL Technology) without feeder cells. After 18–21 days, iPSC colonies were manually picked and expanded in mTeSR-1 maintenance media (STEMCELL Technologies).<sup>27</sup> The iPSCs were routinely passaged every 5–7 days at a ratio of 1:6 with TrypLE (Gibco Life Technologies).

### Validation of the established iPSCs

Morphological images of iPSC colonies were captured with an Olympus CKK53 microscope with an UPlanFL N 4×/0.13na objective lens (Olympus) and Canon digital camera EOS Kiss X10 (Figure S3A). Expression of OCT3/4, SOX2, and NANOG mRNAs was confirmed by real-time quantitative PCR (Figure S3B). Primers sequences are provided in Table S1. Chromosomal G-banding analyses were performed by Chromocentre (Figure S3C). Established iPSCs were differentiated into ectoderm, mesoderm, and endoderm lineages via Pluripotent Stem Cell Functional Identification Kit (R&D Systems) according to the manufacturer's protocol (Figure S3D). In brief, cells were fixed in 4% paraformaldehyde for 20 min at room temperature (RT), washed with PBS, and blocked for 45 min at RT with blocking solution (PBS + 1% BSA + 0.3% Triton-X-100 + 10% normal donkey serum). After blocking, cells were stained with primary antibodies overnight at 4°C. The next day, the cells were washed with PBS and incubated with Anti-Goat Alexa Fluor 488 secondary antibody (ab150129, Abcam) for 1 h at RT. Nuclei were stained with DAPI and the slides were mounted with fluorescent mounting medium (Dako). Images were acquired with a Nikon A1 microscope.

### Chondrogenic differentiation of iPSCs

iPSCs were induced into chondrocytes as described previously.<sup>29</sup> In brief, the iPSCs were initially differentiated into mesendodermal cells in DMEM/F12 (Gibco Life Technologies) with 10 ng/ mL of Wnt-3a (R&D Systems), 10 ng/mL of Activin A (PeproTech), 1% ITS (Gibco Life Technologies), 1% FBS, and 50 units and 50 µg/mL of penicillin and streptomycin. After 3 days, the medium was changed to a chondrogenic medium that consists of DMEM with 1% ITS, 1% FBS, 2 mM L-glutamine,  $1 \times 10^{-4}$  M nonessential amino acids (Wako), 1 mM Na pyruvate (Sigma), 50 µg/mL of ascorbic acid (Wako), 10 ng/mL of BMP2 (Proteintech), 10 ng/mL of TGF- $\beta$ 1 (Proteintech), and 10 ng/mL of GDF5 (PeproTech), 50 units of penicillin, and 50 µg/mL of streptomycin. We added 10 ng/mL of bFGF (Wako) to the chondrogenic medium from day 3 to day 14 to increase the cell proliferation. The medium was changed every 2 days.

### RNA sequencing (RNA-seq)

Total RNA from was extracted with the SV Total RNA Isolation System (Promega) according to the manufacturer's instructions. Total

<sup>(</sup>G and H) Individual 3, born at full term. Short stature with short limbs (G) and oligo-syndactyly of the right hand (H) and the right foot (G). The black box in (G) is used to increase anonymity.

<sup>(</sup>I and J) Lateral view of feet of individuals 1A (I) and 1B (J). Note short feet.

<sup>(</sup>K) Hands of individual 1A. Clinodactyly of the proximal interphalangeal joint of second right digit.

<sup>(</sup>L and M) Dorsal view of feet of individuals 1A (L) and 1B (M). Note short feet and 4-5 syndactyly.



Figure 2. Radiographs of affected individuals
(A) Lower spine of individual 1A with lumbar scoliosis.
(B) Lateral spine of individual 1B showing flattened vertebral bodies and irregular plates.
(C–E) Platyspondyly in individuals 2 (C) and 3 (D and E).

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RNAs were quantified and qualified by Qubit RNA Assay (Thermo Fisher) and TapeStation RNA ScreenTape/BioAnalyzer RNA Pico Kit (Agilent). RNA-seq library preparation was conducted with MGIEasy RNA Directional Library Prep Set V2.0 (MGI Tech). The libraries were fed into a DNBSEQ-G400 sequencer (MGI Tech) according to the manufacturer's instructions, with paired end reads and mean read lengths of 150 bp. Read quality was assessed and adapters were trimmed from the reads via fastp v.0.20.0. RNAseq yielded about 40 million reads per sample.

### Differential gene expression analysis of RNA-seq data

The reads from RNA-seq were aligned to the human genome (GRCh38) with HISAT2 v 2.2.0. Aligned reads were quantified with featureCounts 2.6.0. Transcripts per million (TPM) counts was calculated by Python package bioinfokit v2.0.8. The volcano plot and the ridgeline plot were created with ExpressAnalyst. The heatmaps were created by GraphPad Prism 8.0. Enrichment analysis was performed by ExpressAnalyst.

### Alcian blue staining

Cells were fixed with 95% methanol for 20 min and stained with 1% Alcian blue 8GX (Sigma) in 0.1 M HCL overnight.<sup>30</sup> We washed the stained cells aggregated in plates at least three times with deionized water to guarantee only specific staining was left and we scanned these cells with a light microscope (Olympus CK40).

### Statistical analysis

Continuous variables were reported as mean  $\pm$  SD and compared with a two-tailed Student's t test or a one-way ANOVA followed by a post-hoc Tukey's multiple-comparison test as specified in the figure legends. p values < 0.05 were considered to be significant.

### Results

# A phenotypic dichotomy is identified in individuals with *ERI1* variants

*ERI1* variants were identified as candidates for the severe skeletal dysplasia by ES analysis independently for families 1, 2, and 3. It was identified as a candidate for the distal limb anomalies independently by ES analysis for families 5 and 6. Investigators were in contact through GeneMatcher. Family 4 was identified by a search of the Baylor Genetics Laboratories exome variant database for individuals with two *ERI1* potentially deleterious variants, and family 7

was identified by a search of the 100,000 Genomes Project variant database. Six missense variants, three nonsense variants, and one splice-site variant of *ERI1* were identified in these affected individuals (Table 1). Homozygosity or compound heterozygosity of these variants was confirmed in the individuals except individual 4, who harbored p.Ser21\* and p.Pro155Leu, as the parents were not available. The frequency of these variants was less than 0.0001 (c.464C>T [p.Pro155Leu], c.893A>G [p.Asp298Gly]) or not present (the others) in gnomAD and ExAC.

Although various digital abnormalities existed in all affected individuals (Figures 1E–1N and 2Q–2W), SEMD was only present in those with missense *ERI1* variants (families 1–4) (Figures 2A–2P, Table 1). Additional clinical manifestations, including cardiac anomalies, developmental delay, and hydronephrosis, were found in some of the individuals (Table 1). Clinical summaries are provided in the supplemental note and pedigrees in Figure S1. A search of known monogenic skeletal disorders, including disease entities registered in the latest version of the international nosology and classification of genetic skeletal disorders,<sup>31</sup> did not find any reports matching the characteristic SEMD that was observed in this study. We tentatively suggest the name SEMD, Guo-Campeau type for this phenotype.

In contrast, three individuals with bi-allelic *ERI1* null mutations (families 5–7) showed a much milder skeletal phenotype, which affects the patterning of digital bones but without any chondrodysplastic feature such as SEMD (Table 1). The prominent clinical manifestation of these individuals was intellectual disability (Table 1), consistent with the previous individual with homozygous 284 kb deletion reported by the Mégarbané's group.<sup>10</sup>

# The *ERI1* variants exhibit loss of 3'-to-5' exoribonuclease activity

ERI1 includes two functional domains: the amino-terminal SAP (SAF-A/B, Acinus, and PIAS) domain and the carboxy-terminal 3' exonuclease domain (Figure 3A). The structure of the exonuclease domain complexed with riboadenine-5'-monophosphate (rAMP) is determined in the presence of Mg<sup>2+</sup> ions (Figure 3B).<sup>32</sup> The six missense variants, p.Asp134Gly, p.Glu150Asp, p.Pro155Leu, p.Asp298Gly, p.Asp298Ala, and p.Ser299Pro, identified in our study are all located in the

<sup>(</sup>F–M) Long tubular bones showing epi-metaphyseal dysplasia characterized by irregular epiphyses and frayed metaphyses in individuals 1A (F), 1B (G), 2 (H and I), and 3 (J–M). The right upper limb of individual 3 shows ulnar hypoplasia and radial bowing with elbow dislocation (J).

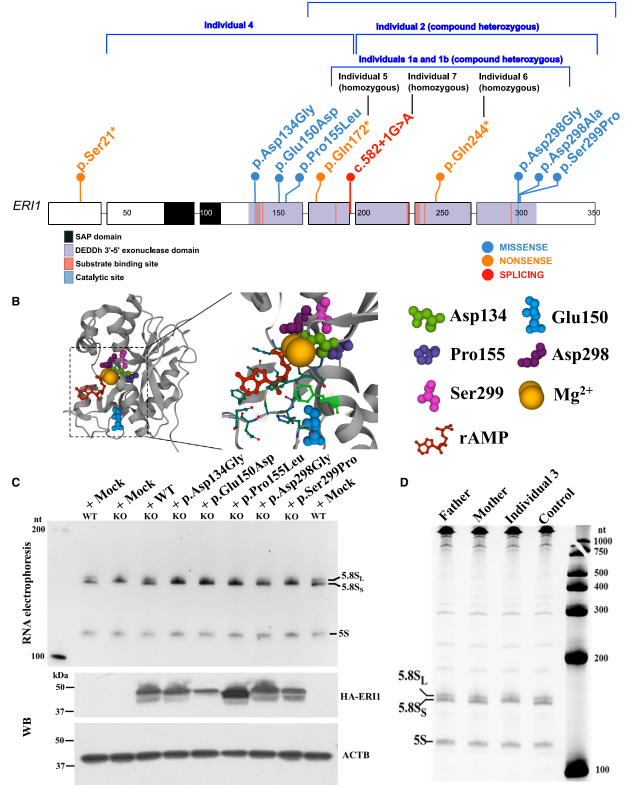
<sup>(</sup>N–P) Hypoplasia of pelvis and epi-metaphyseal dysplasia of femurs in individuals 1B (N), 2 (O), and 3 (P). Note right coxa vara and left coxa valga (N), iliac hypoplasia and delayed pubic ossification (O and P).

<sup>(</sup>Q) Right hand of individual 1A. Note single thumb phalanx, ulnar deviation of the second finger at proximal interphalangeal joint, and narrow metacarpal bones.

<sup>(</sup>R) Left hand of individual 2 showing brachy-syndactyly of the third and fourth fingers.

<sup>(</sup>S–V) Hands and feet of individual 3. The left hand (S) shows brachydactyly with hypoplasia of all middle phalanges, the first proximal and distal phalanges, and the second and third distal phalanges as well as ulnar clinodactyly of the second and third fingers and radial clinodactyly of the fourth and fifth fingers. The proximal end of the proximal phalanges and distal ends of the metacarpals are conspicuously cupped. The right hand shows oligo-syndactyly (split hand) (T). Both feet show postaxial syndactyly and the third and fourth metatarsal bases may be fused (U and V). The right fourth metatarsal is hypoplastic (V).

<sup>(</sup>W) Right foot of individual 7. Note absence of the fifth ray, proximal fusion of the third and fourth metatarsal bones, and widened fourth metatarsus. The black boxes in (J), (L), and (U) mask arrows previously used in clinical discussions.



#### Figure 3. Functional analysis of ERI1 pathogenic variants

(A) Schematic overview of ERI1 showing the position of the variants identified in the individuals. ERI1 includes two functional domains: amino-terminal SAP (SAF-A/B, Acinus, and PIAS) domain and carboxy-terminal 3' exonuclease domain.
(B) 3D structure of ERI1 with annotated residues Asp134, Glu150, Pro155, Asp298, and Ser299 generated via UniProKB (https://www.uniprot.org/uniprot/Q8IV48). PDB: 1ZBU. rAMP, riboadenine-5'-monophosphate.

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carboxy-terminal 3' exonuclease domain of ERI1 (Figure 3A). A 3D structure of ERI1 indicates that the residues corresponding to these missense variants are parts of the active site of the nuclease domain and are required to coordinate the two  $Mg^{2+}$  ions (Figure 3B), thus suggesting their key roles for ERI1 enzyme activity. c.582+1G>A was a canonical splicedisrupting variant. The location of the nonsense variants p.Ser21\*, p.Gln172\*, and p.Gln244\* suggests that all of them are subjected to nonsense-mediated mRNA decay.

Eri1 is a 3'-to-5' exoribonuclease that catalyzes 5.8S rRNA processing.<sup>7,9</sup> To study a potential deleterious effect of these missense ERI1 variants, we generated an ERI1 KO HeLa cell line (Figure S2) and demonstrated that ERI1 was required for proper 5.8 rRNA 3' end formation also in human cells (Figures 3C and S2F), consistent with the previous findings in mice, worms, and fission yeast.<sup>7,9</sup> Then we attempted to rescue defective 5.8S rRNA processing by generating ERI1 from the WT construct and the missense variant constructs in KO HeLa cells. While WT ERI1 efficiently catalyzed conversion of the abnormal 5.8S rRNA to its normal size, all of the variant ERI1s failed to do so (Figure 3C). Consistently, electrophoresis of total RNA extracted from the family 3-derived lymphoblastoid cells showed that the affected individual, but not his parents or an unrelated unaffected individual, exhibited the same 5.8S rRNA processing defect as the KO HeLa cells (Figure 3D). Thus, both in vitro and in vivo evidence indicates that the ERI1 missense variants cause a loss of function of the 3'-to-5' exoribonuclease activity of ERI1.

### Effect of ERI1 deficiency on abundance of replicationdependent histone mRNAs and miRNAs

Besides 5.8S rRNA processing, Eri1 regulates the abundance of histone mRNAs and miRNAs in mouse cells.<sup>33</sup> In mouse embryonic fibroblasts, loss of Eri1 leads to excessive accumulation of replication-dependent histone mRNAs.<sup>8</sup> To assess the corresponding function in human cells, we analyzed histone mRNA levels in the mother- and affected-individual-derived iPSCs from family 3 by using RNA-seq data. Indeed, the replication-dependent histone mRNAs showed an overall increased abundance in the affected-individual-derived iPSCs compared to the replication-independent histone mRNAs (Figure 4A), in accordance with the previous findings in mouse cells.<sup>8</sup> This effect was also evident in the ERI1 KO HeLa cells and affected-individual-derived iPSCs and fibroblasts treated with hydroxyurea (Figures 4B-4D), which induces degradation of replication-dependent histone mRNAs similar to their physiologic regulation at the end of S phase.<sup>8</sup> These findings confirm the regulatory role of ERI1 in this

class of mRNAs as shown in mouse cells<sup>8</sup> and indicate a decreased degradation of replication-dependent histone mRNAs in affected individuals.

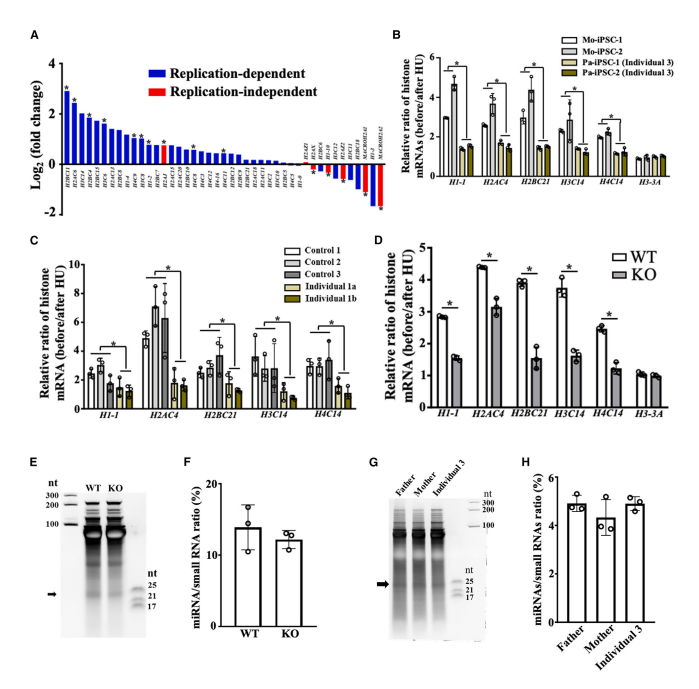
Additionally, conditional deletion of *Eri1* in mouse natural killer (NK) and T cells displayed a global, sequence-independent increase in miRNA abundance.<sup>6</sup> We electrophoresed total small RNAs shorter than 300 nucleotides (nts) extracted from WT and *ERI1* KO HeLa cells and affected-individual-derived lymphoblastoid cell lines to evaluate the proportion of miRNAs (Figures 4E and 4G). We enhanced the visualization of small RNAs by using SYBR Gold Nucleic Acid Gel Stain.<sup>34</sup> As a result, a smear of bands corresponding to single-strand RNAs (ssRNAs) between 21 and 25 nts was detected; however, the overall abundance of these small RNA species did not change in *ERI1* KO HeLa cells or affected-individual-derived lymphoblastoid cells (Figures 4F and 4H).

# Impaired chondrogenesis in affected-individual-derived iPSCs

To examine possible roles of ERI1 in chondrogenesis that could then explain the etiology of the individuals' skeletal phenotypes, we established and verified four iPSC lines by using affected-individual-derived lymphoblastoid cell lines (Figure S3), which were generated from peripheral blood lymphocytes of an affected individual and his mother from family 3. The iPSC lines were differentiated toward chondrocytes as described previously.<sup>29</sup> While Alcian blue (AB) staining of sulfated glycosaminoglycan accumulated in the mother-derived lines at day 21, significantly decreased AB staining was found in the affect-individualderived lines (Figures 5A and 5B). The expression of the chondrocyte marker COL2A1 was also decreased in the affected-individual-derived lines, as compared to its expression in the mother-derived lines (Figure 5C). We then subjected the differentiating iPSCs at day 8 (which correspond to mesodermal cells<sup>29</sup>) to RNA-seq. We identified 634 differentially expressed genes (Figure 5D). A gene set enrichment analysis identified several Gene Ontology (GO) terms, including "skeletal system development" and "pattern specification process" (Figure 5E), in which the genes were mostly downregulated in the affected-individual-derived cells. The downregulated genes included DLX5, DLX6, PKDCC, BMP4, SUFU, ROR2, TBX3, TRAPPC2B, and GDF11, which are associated with skeletal dysplasias,<sup>35–41</sup> thereby uncovering disease-relevant changes in mesodermal differentiation of cells from affected individuals (Figure 5F). These skeletal dysplasias have defective development in the short tubular bones, such as split hand/foot malformation (MIM: 183600) and

<sup>(</sup>C) The upper panel shows ethidium bromide-stained RNA from the wild type (WT) and *ERI1* knockout (KO) HeLa cells. Nt, nucleotide. WT, p.Asp134Gly, p.Glu150Asp, p.Pro155Leu, p.Asp298Gly, and p.Ser299Pro ERI1 were produced in the KO cells to rescue the 3'-to-5' exoribonuclease activity of ERI1 in catalyzing 5.8S ribosomal RNA (rRNA) processing. The lower panel shows western blot of whole-cell lysates.

<sup>(</sup>D) Ethidium bromide-stained RNA from the lymphoblastoid cell lines of the affected individual 3, his parents, and a healthy control.  $5.8S_{L}$  long form of 5.8S rRNA;  $5.8S_{S}$ , short form of 5.8S rRNA.



#### Figure 4. Effect of ERI1 deficiency on histone mRNAs and miRNAs

(A) Histone mRNA levels of the family 3-derived iPSCs in the process of chondrogenic differentiation (day 8). Both replication-dependent and replication-independent histone genes are ranked according to the fold change (FC) of the differentially expressed genes in MoiPSC-1 versus Pa-iPSC-1. \*p < 0.05. n = 3 biologically independent samples. Statistical significance was assessed via two-sided t test. p value was adjusted by false discovery rate (FDR).

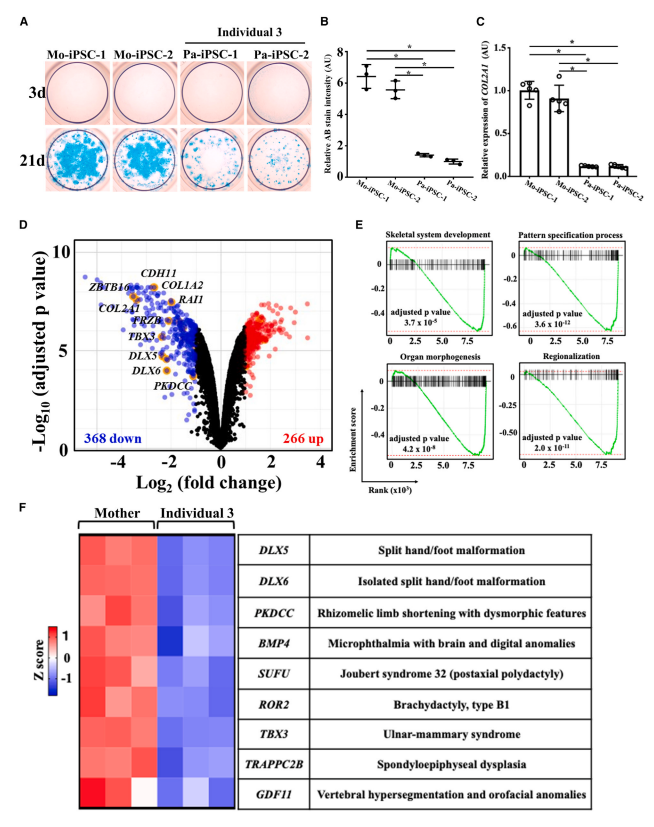
(B–D) qPCR for histone mRNAs before and after hydroxyurea (HU) treatment for 45 min. All markers are replication-dependent histone genes except *H3-3A*.

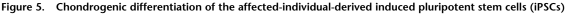
(B) Family 3-derived iPSCs.

(C) Fibroblasts from three healthy control individuals and the two affected individuals in family 1.

(D) Wild type (WT) and knock-out (KO) HeLa cells. The data in (B)–(D) indicate mean  $\pm$  SD. n = 3 biological replicates. Statistical test was assessed via multiple t tests. \*p < 0.05.

(E–H) SYBR Gold-stained small RNAs from the wild type (WT) and *ERI1* knockout (KO) HeLa cells (E) and family 3-derived lymphoblastoid cells (G). nt, nucleotide. The arrows show single-strand RNAs between 21 and 25 nt, corresponding to the abundance of miRNAs. Quantification of the miRNA to total small RNA (<300 nt) ratio in HeLa cells (F) and lymphoblastoid cells (H). The data indicate mean  $\pm$  SD. p = 0.44 and 0.33. n = 3 biological replicates. Statistical significance was assessed via two-sided t test and one-way ANOVA.





(A) Alcian blue (AB) staining for the chondrocytic nodules induced from the healthy mother iPSCs (Mo-iPSC-1 and -2) and affected-individual-derived iPSCs (Pa-iPSC-1 and -2) in family 3.

(B) Quantification of AB staining; n = 3 biologically independent samples. \*p < 0.05. Error bars represent SD.

(C) qPCR for *COL2A1*; n = 5 biologically independent samples. \*p < 0.05. AU, arbitrary unit. The data in (B) and (C) indicate mean  $\pm$  SD. Statistical test was assessed via one-way ANOVA.

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brachydactyly, type B1 (MIM: 113000), and in the long tubular bones and spine, such as spondyloepiphyseal dysplasia (Figure 5F), which are similar to the skeletal phenotypes of affected individuals described here. These results indicate that affected-individual-derived iPSCs with bi-allelic missense variants are impaired in their ability to undergo chondrogenesis, suggesting that defects in both digital patterning and chondrogenesis are caused by missense variants.

# *Eri1* KO mice partially recapitulate skeletal dysplasia caused by human *ERI1* pathogenic variants

Eri1 KO mice on the C57BL/6 inbred strain background show a growth defect with reduced body size and neonatal mortality.<sup>7</sup> Neonatal lethality was rescued by breeding Eri1 KO mice on outbred strain backgrounds.<sup>7</sup> To test potential phenotypic similarities between the affected individuals and the KO mice, we evaluated the morphological changes of the Eri1 KO mice skeleton on an NMRI × C57BL/6 mixed background. The Eri1 KO mice showed brachydactyly (Figures 6A and 6B), which was also characteristic for the affected individuals with ERI1 variants (Figures 2Q-2W). The examination of X-rays and histological stain did not show obvious change in the epimetaphyseal regions of the long tubular bones of the Eri1 KO mice (Figure S4). Although the Eri1 KO mice showed platyspondyly (Figures 6C and 6D), which was also characteristic for the affected individuals with ERI1 missense variants (Figures 2A-2E), such a phenotype was mild in the KO mice, perhaps being secondary to the reduced body size. Therefore, the skeletal phenotypes of Eri1 KO mice were more similar to the affected individuals with null variants than those with missense variants.

### Discussion

In this study, we describe a type of SEMD by characterizing the phenotypes caused by different types of bi-allelic *ERI1* variants. The SEMD was present in the five individuals with at least one missense variant (Table 1). In contrast, three individuals with *ERI1* null mutations and the *Eri1* KO mice showed a much milder skeletal phenotype without any evidence for SEMD, consistent with the two individuals reported previously, who had homozygous a 284 kb deletion and p.Lys118\*. Notably, of the five individuals with SEMD, three died within 2 years after birth, suggesting missense variants lead to a poor prognosis.

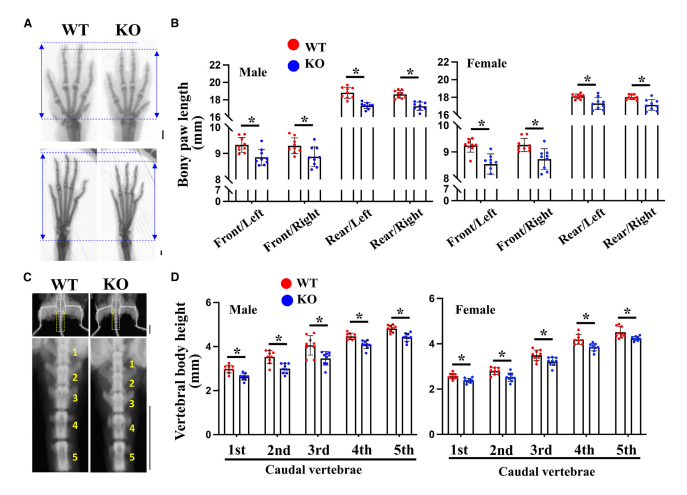
The phenomenon that more severe clinical phenotypes are associated with missense variants rather than null variants is frequently observed in some dominant disorders, such as osteogenesis imperfecta caused by COL1A1 and COL1A2 mutations,<sup>42</sup> but not reported in recessive disorders. All missense variants in affected individuals are located in the carboxy-terminal 3' exonuclease domain without disrupting the sequence of the amino-terminal SAP domain, which is an RNA-binding motif. These exonuclease-dead proteins may compete for the target RNA molecules with other exonucleases that have functional redundancy with ERI1, staying bound to those RNA molecules and thus leading to a more severe phenotype in skeleton (Figure 7). Alternatively, these exonuclease-dead proteins may impair the chondrogenesis by an additional mechanism (Figure 7). The genotype-phenotype association needs to be verified further by finding more affected individuals and comparing the phenotypic difference between the KO mice and the knockin mice to be generated with the individual-specific missense variants.

Eri1 is a 3'-to-5' exoribonuclease, whose functions generally show evolutionary conservation in the processing of 5.8S rRNA and histone mRNAs and species-specific differences in the regulation of small RNA pathways that include endogenous siRNAs and miRNAs.9,33,43 The 5.8S rRNA 3' end forms a duplex reminiscent of pre-miRNAs by base pairing with the 5' end of the 28S rRNA, which leaves a 3' ssRNA overhang processed by Eri1 into a duplex with 1-2 nt 3' overhang.<sup>7,9</sup> Our study demonstrated that ERI1 processes 5.8S rRNA in humans as it does in other species. Although the biological significance of the ERI1-mediated 5.8S rRNA 3' trimming remains unknown, the biological process is conserved from yeast to humans, suggesting its importance in ribosome biogenesis, a complex endeavor that involves coordination of three RNA polymerases, approximately 80 ribosomal proteins (RPs), 70 small nucleolar RNAs, and several rRNAs.<sup>44</sup>

Variants in ribosomal proteins and regulators of ribosome biogenesis lead to ribosomopathies, a group of disorders that show developmental defects in craniofacial bones and long and short tubular bones, causing skeletal dysplasias such as Treacher Collins syndrome 1 (MIM: 154500), postaxial acrofacial dysostosis, Shwachman-Diamond syndrome 1 (MIM: 260400), Roberts syndrome, and cartilage hair hypoplasia (MIM: 250250).<sup>45–49</sup> The skeletal phenotypes of these disorders include spondylo-metaphyseal dysplasia and digit deformities, showing similarities with affected individuals reported here. Ribosomopathies are also characterized by hematologic phenotypes such as macrocytic anemia in Diamond-Blackfan anemia 1 (MIM: 105650) and congenital neutropenia in Shwachman-Diamond-like syndrome.<sup>50–52</sup> Macrocytic anemia and

(F) Heatmap of DEGs associated causally with human skeletal dysplasia.

<sup>(</sup>D) Volcano plot of RNA-seq data for the differentiating iPSCs at day 8 showing differentially expressed genes (DEGs) in Mo-iPSC-1 versus Pa-iPSC-1 (n = 3 biologically independent samples) with respect to fold change (FC) and significance (adjusted p value). Each dot represents an individual gene. Blue dots represent significant downregulated DEGs (FC < 0.5) and red dots represent significant up-regulated DEGs (FC > 2). Statistical significance was assessed via moderated t test in limma. p value is adjusted by false discovery rate. (E) Gene set enrichment analysis for RNA-seq data in (D). The genes belonging to the corresponding GO terms are enriched at the bottom of the entire ranked list of genes.



### Figure 6. Skeletal dysplasia of Eri1 knockout (KO) mice

(A) X-ray images of paws of the wild type (WT) and Eri1 KO mice at 16 weeks. Scale bar, 1 mm.

(B) Quantitative analysis of the paw length shown in (A) (n = 9 mice). \*p < 0.05. The above quantitative data indicate mean  $\pm$  SD and statistical significance was assessed via two-sided t test. Error bars represent SD.

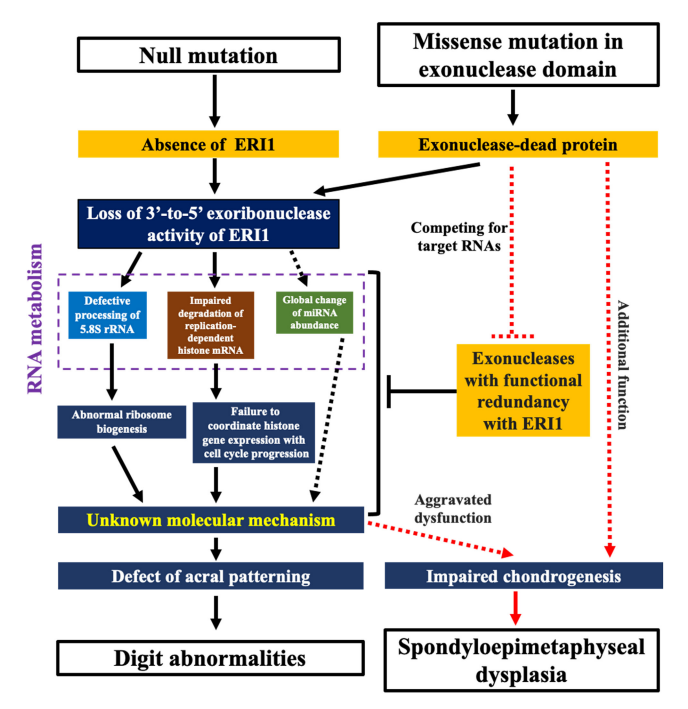
(C) X-ray images of the WT and *Eri1* KO mice at 16 weeks. The areas highlighted by the dotted box in the upper panels are magnified in the lower panels. Scale bar, 1 cm.

(D) Measurement of the height of the caudal vertebral bodies shown in (C) (n = 9 mice). \*p < 0.05.

congenital neutropenia were also evident in individual 3 and the *Eri1* KO mice (Figures S5 and S6). Thus, both the biological function of ERI1 and the clinical manifestations suggest existence of a ribosomopathy in the individuals reported here. The known genes associated with ribosomopathies cover functions in methylation and cleavage of 47S precursor rRNA, 18S rRNA maturation, and assembly of ribosomal subunits.<sup>53</sup> The defective processing of 5.8S rRNA in our cohort would provide a link that associates human disease with ribosome biogenesis, thus enriching the present list of ribosomopathies.

Eri1 participates in fast degradation of replication-dependent histone mRNAs in mice by binding to the oligouridylated histone mRNA 3' end and trimming two nucleotides at the end of S phase, similar to its role in 5.8S rRNA maturation.<sup>8</sup> Marked accumulation of replication-dependent histone mRNAs were observed in our affected-individualderived cells and ERI1 KO HeLa cells as found in the *Eri1* KO mouse cells, indicating the function of histone mRNA exonuclease of ERI1 is also conserved in humans. High levels of histone mRNAs are closely linked to DNA replication so that numerous histones are produced in S phase to accommodate DNA packaging into nucleosomes. Histone mRNA abundance then falls rapidly in G2 by fast degradation. Although the biological effects of excessive replication-dependent histone mRNAs remain unknown in mammals, failure to coordinate histone gene expression with cell-cycle progression results in genomic instability and cell-cycle arrest in yeast.<sup>54</sup> Our study suggests that the defective histone mRNA decay may disrupt skeletal development singly or in combination with the defective 5.8S rRNA maturation in the affected individuals.

Previous studies using *Eri1* KO mice reported that Eri1 loss causes a global, sequence-independent increase in mature miRNA abundance in NK and T cells.<sup>6</sup> Unexpectedly, we did not detect a global increase in the abundance of 21–25 nt ssRNAs in *ERI1* KO HeLa cells and affected-individual-derived lymphoblastoid cells. *Eri1* is highly expressed in mouse spleen and thymus compared to other tissues.<sup>7</sup> NK and T lymphocytes upregulate *Eri1* expression to high levels



# Figure 7. Hypothesis of the pathological mechanism underlying the phenotypic dichotomy between null and missense variants of *ERI1*

Both null and missense mutations of *ERI1* cause loss of 3'-to-5' exoribonuclease activity of ERI1, leading to defective processing of 5.8S rRNA and impaired degradation of replication-dependent histone mRNA in humans, which interrupt diverse cellular processes including ribosome biogenesis and coordination of histone gene expression with cell cycle progression. These defects singly or jointly alter gene expression profiling that regulates acral patterning, thus inducing digit abnormalities. *ERI1* missense mutations also produce exonuclease-dead proteins, which may compete with other exonucleases having functional redundancy with ERI1, thus aggravating the dysfunction of ERI1-mediated RNA metabolism. The aggravated dysfunction or additional unknown effects lead to impaired chondrogenesis that causes spondyloepimetaphyseal dysplasia, finally contributing to the phenotypic dichotomy between null and missense variants of *ERI1*. Solid arrow, causal connection; dashed arrow, hypothetical connection; red line, additional effects of exonuclease-dead ERI1 to the loss of the exoribonuclease activity.

upon activation, thus suggesting the Eri1 effects on miRNA turnover may be cell/context-dependent.<sup>6</sup> Alternatively, the regulatory role of ERI1 in miRNA homeostasis may not be completely conserved between mice and humans.

The multiple RNA metabolic pathways regulated by Eri1 make the deeply conserved exoribonuclease play an extremely complex role in the crosstalk and co-regulation of diverse cellular processes. Our study establishes a causal

link between ERI1 deficiency and human skeletal dysplasia and reveals the evolutionarily conserved functions of ERI1 in RNA metabolic pathways, which associate 5.8S rRNA maturation and histone mRNA decay with the regulation of bone morphogenesis (Figure 7), thus leading to further exploration of the roles of ERI1-mediated RNA metabolism in skeletal development. Our study reveals a phenotypic dichotomy between null and missense variants of *ERI1*, thus serving as a precedent for a nuanced form of mechanistic heterogeneity for recessive conditions in humans (Figure 7).

### Data and code availability

The RNA-seq data have been deposited in NCBI GEO with the accession code GEO: GSE206412. The individuals' genomic data obtained by exome or genome sequencing are not publicly available because study participants did not give full consent for releasing data publicly. These data can be accessed under the condition that a joint research plan is made by the researchers and approved by the ethics committees of the participating institutions. All the other data supporting the findings of this study are included within the article or the supplemental information.

### Supplemental information

Supplemental information can be found online at https://doi.org/ 10.1016/j.ajhg.2023.06.001.

### Author contributions

The study was designed by L.G., S.I., and P.M.C. The individuals were recruited by N.N., H.K., J.C.S., N.D.N., S.B., A.J., I.H., T.B., C.B., F.A., R.M., G.C., B.W., A.M., T.Y., G.N., J.A.R., S.I., and P.M.C. The disease was evaluated by L.G., T.Y., G.N., S.I., and P.M.C. The experiments were performed or supervised by L.G., S.S., J.R., H.K., J.Y.X., N.R., Z.W., Z.J.W., H.O., M.F.T., N. Matsumoto, N. Miyake, M.K.A., S.M., G.Y., B.W., M.H.d.A., and V.H. The data were analyzed by L.G., S.S., J.Y.X., J.W., and Y.M.Z. The draft was written by L.G., S.I., and P.M.C. All authors contributed to revising the manuscript.

### **Declaration of interests**

The Department of Molecular and Human Genetics at Baylor College of Medicine receives revenue from clinical genetic testing completed at Baylor Genetics Laboratories.

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#### Web resources

bioinfokit, https://pypi.org/project/bioinfokit/ ClinVar, https://www.ncbi.nlm.nih.gov/clinvar/ dbSNP, https://www.ncbi.nlm.nih.gov/projects/SNP/ E-CRISP, http://www.e-crisp.org/E-CRISP/ ExpressAnalyst, https://www.expressanalyst.ca/ fastp https://github.com/OpenGene/fastp featureCounts, https://subread.sourceforge.net/ GeneMatcher https://genematcher.org/ gnomAD population database, https://gnomad.broadinstitute.org/ HISAT2, http://daehwankimlab.github.io/hisat2/

Human Gene Mutation Database, https://portal.biobase-inter national.com/hgmd/pro/start.php

Human Genome Variation Society guidelines, http://www.hgvs.org limma https://bioconductor.org/packages/release/bioc/html/lim ma.html

OMIM, https://omim.org/

UniProKB, https://www.uniprot.org/uniprot/

### References

- 1. Zuo, Y., and Deutscher, M.P. (2001). Exoribonuclease superfamilies: Structural analysis and phylogenetic distribution. Nucleic Acids Res. *29*, 1017–1026.
- Simmer, F., Tijsterman, M., Parrish, S., Koushika, S.P., Nonet, M.L., Fire, A., Ahringer, J., and Plasterk, R.H.A. (2002). Loss of the putative RNA-directed RNA polymerase RRF-3 makes C. Elegans hypersensitive to RNAi. Curr. Biol. *12*, 1317–1319.
- **3.** Gu, W., Shirayama, M., Conte, D., Vasale, J., Batista, P.J., Claycomb, J.M., Moresco, J.J., Youngman, E.M., Keys, J., Stoltz, M.J., et al. (2009). Distinct Argonaute-Mediated 22G-RNA Pathways Direct Genome Surveillance in the C. elegans Germline. Mol. Cell *36*, 231–244.
- 4. Han, T., Manoharan, A.P., Harkins, T.T., Bouffard, P., Fitzpatrick, C., Chu, D.S., Thierry-Mieg, D., Thierry-Mieg, J., and Kim, J.K. (2009). 26G endo-siRNAs regulate spermatogenic and zygotic gene expression in Caenorhabditis elegans. Proc.-Natl.Acad.Sci.USA *106*, 18674–18679.
- Bühler, M., Verdel, A., and Moazed, D. (2006). Tethering RITS to a Nascent Transcript Initiates RNAi- and Heterochromatin-Dependent Gene Silencing. Cell *125*, 873–886.
- 6. Thomas, M.F., Abdul-Wajid, S., Panduro, M., Babiarz, J.E., Rajaram, M., Woodruff, P., Lanier, L.L., Heissmeyer, V., and Ansel, K.M. (2012). Eri1 regulates microRNA homeostasis and mouse lymphocyte development and antiviral function. Blood *120*, 130–142.
- Ansel, K.M., Pastor, W.A., Rath, N., Lapan, A.D., Glasmacher, E., Wolf, C., Smith, L.C., Papadopoulou, N., Lamperti, E.D., Tahiliani, M., et al. (2008). Mouse Eri1 interacts with the ribosome and catalyzes 5.8S rRNA processing. Nat. Struct. Mol. Biol. 15, 523–530.
- Hoefig, K.P., Rath, N., Heinz, G.A., Wolf, C., Dameris, J., Schepers, A., Kremmer, E., Ansel, K.M., and Heissmeyer, V. (2013). Eri1 degrades the stem-loop of oligouridylated histone mRNAs to induce replication-dependent decay. Nat. Struct. Mol. Biol. 20, 73–81.
- **9.** Gabel, H.W., and Ruvkun, G. (2008). The exonuclease ERI-1 has a conserved dual role in 5.8S rRNA processing and RNAi. Nat. Struct. Mol. Biol. *15*, 531–533.
- Choucair, N., Rajab, M., Mégarbané, A., and Chouery, E. (2017). Homozygous microdeletion of the *ERI1* and *MFHAS1* genes in a patient with intellectual disability, limb abnormalities, and cardiac malformation. Am. J. Med. Genet. *173*, 1955–1960.
- **11.** Hoxha, V., and Aliu, E. (2023). ERI1: A case report of an autosomal recessive syndrome associated with developmental delay and distal limb abnormalities. Am. J. Med. Genet. *191*, 64–69.
- Ravera, S., Dufour, C., Cesaro, S., Bottega, R., Faleschini, M., Cuccarolo, P., Corsolini, F., Usai, C., Columbaro, M., Cipolli, M., et al. (2016). Evaluation of energy metabolism and

calcium homeostasis in cells affected by Shwachman-Diamond syndrome. Sci. Rep. *6*, 25441.

- **13.** (2017). Inflammatory Signature, Oxidative Stress, and DNA Damage Response in DBA Pathogenesis. Blood *130*, 2452.
- 14. Zambetti, N.A., Ping, Z., Chen, S., Kenswil, K.J.G., Mylona, M.A., Sanders, M.A., Hoogenboezem, R.M., Bindels, E.M.J., Adisty, M.N., Van Strien, P.M.H., et al. (2016). Mesenchymal Inflammation Drives Genotoxic Stress in Hematopoietic Stem Cells and Predicts Disease Evolution in Human Pre-leukemia. Cell Stem Cell 19, 613–627.
- Hermanns, P., Bertuch, A.A., Bertin, T.K., Dawson, B., Schmitt, M.E., Shaw, C., Zabel, B., and Lee, B. (2005). Consequences of mutations in the non-coding RMRP RNA in cartilage-hair hypoplasia. Hum. Mol. Genet. *14*, 3723–3740.
- Brunet, T., Jech, R., Brugger, M., Kovacs, R., Alhaddad, B., Leszinski, G., Riedhammer, K.M., Westphal, D.S., Mahle, I., Mayerhanser, K., et al. (2021). De novo variants in neurodevelopmental disorders-experiences from a tertiary care center. Clin. Genet. *100*, 14–28.
- 17. Yang, Y., Muzny, D.M., Xia, F., Niu, Z., Person, R., Ding, Y., Ward, P., Braxton, A., Wang, M., Buhay, C., et al. (2014). Molecular findings among patients referred for clinical wholeexome sequencing. JAMA *312*, 1870–1879.
- Xue, J.Y., Grigelioniene, G., Wang, Z., Nishimura, G., Iida, A., Matsumoto, N., Tham, E., Miyake, N., Ikegawa, S., and Guo, L. (2022). <scp> *SLC4A2* </scp> Deficiency Causes a New Type of Osteopetrosis. J. Bone Miner. Res. *37*, 226–235.
- Guo, L., Iida, A., Bhavani, G.S., Gowrishankar, K., Wang, Z., Xue, J.Y., Wang, J., Miyake, N., Matsumoto, N., Hasegawa, T., et al. (2021). Deficiency of TMEM53 causes a previously unknown sclerosing bone disorder by dysregulation of BMP-SMAD signaling. Nat. Commun. *12*, 2046–2113.
- Guo, L., Elcioglu, N.H., Karalar, O.K., Topkar, M.O., Wang, Z., Sakamoto, Y., Matsumoto, N., Miyake, N., Nishimura, G., and Ikegawa, S. (2018). Dysosteosclerosis is also caused by TNFRSF11A mutation. J. Hum. Genet. 63, 769–774.
- 21. Guo, L., Bertola, D.R., Takanohashi, A., Saito, A., Segawa, Y., Yokota, T., Ishibashi, S., Nishida, Y., Yamamoto, G.L., Franco, J.F.d.S., et al. (2019). Bi-allelic CSF1R Mutations Cause Skeletal Dysplasia of Dysosteosclerosis-Pyle Disease Spectrum and Degenerative Encephalopathy with Brain Malformation. Am. J. Hum. Genet. 104, 925–935.
- 22. Stavropoulos, D.J., Merico, D., Jobling, R., Bowdin, S., Monfared, N., Thiruvahindrapuram, B., Nalpathamkalam, T., Pellecchia, G., Yuen, R.K.C., Szego, M.J., et al. (2016). Whole Genome Sequencing Expands Diagnostic Utility and Improves Clinical Management in Pediatric Medicine. NPJ Genom. Med. 1, 15012.
- 23. Cong, L., Ran, F.A., Cox, D., Lin, S., Barretto, R., Habib, N., Hsu, P.D., Wu, X., Jiang, W., Marraffini, L.A., and Zhang, F. (2013). Multiplex genome engineering using CRISPR/Cas systems. Science *339*, 819–823.
- 24. Humbert, J., Salian, S., Makrythanasis, P., Lemire, G., Rousseau, J., Ehresmann, S., Garcia, T., Alasiri, R., Bottani, A., Hanquinet, S., Beaver, E., Heeley, J., Smith, A.C.M., Berger, S.I., Antonarakis, S.E., Yang, X.J., Côté, J., and Campeau, P.M. (2020). De Novo KAT5 Variants Cause a Syndrome with Recognizable Facial Dysmorphisms, Cerebellar Atrophy, Sleep Disturbance, and Epilepsy. Am. J. Hum. Genet. *107*, 564–574.
- 25. Fuchs, H., Gailus-Durner, V., Adler, T., Pimentel, J.A.A., Becker, L., Bolle, I., Brielmeier, M., Calzada- Wack, J., Dalke, C., Ehrhardt, N., et al. (2009). The German Mouse Clinic: A Platform

for Systemic Phenotype Analysis of Mouse Models. Curr. Pharm. Biotechnol. *10*, 236–243.

- **26.** Fukushima, Y., Ohashi, H., Wakui, K., Nishida, T., and Oh-Ishi, T. (1992). A rapid method for starting a culture for the establishment of Epstein-Barr virus-transformed human lymphoblastoid cell lines. J. Hum. Genet. *37*, 149–150.
- 27. Kumar, S., Curran, J.E., Espinosa, E.C., Glahn, D.C., and Blangero, J. (2020). Highly efficient induced pluripotent stem cell reprogramming of cryopreserved lymphoblastoid cell lines. J. Biol. Methods 7, e124.
- 28. Okita, K., Matsumura, Y., Sato, Y., Okada, A., Morizane, A., Okamoto, S., Hong, H., Nakagawa, M., Tanabe, K., Tezuka, K.I., et al. (2011). A more efficient method to generate integration-free human iPS cells. Nat. Methods *8*, 409–412.
- 29. Yamashita, A., Morioka, M., Yahara, Y., Okada, M., Kobayashi, T., Kuriyama, S., Matsuda, S., and Tsumaki, N. (2015). Generation of scaffoldless hyaline cartilaginous tissue from human iPSCs. Stem Cell Rep. *4*, 404–418.
- 30. Shukunami, C., Shigeno, C., Atsumi, T., Ishizeki, K., Suzuki, F., and Hiraki, Y. (1996). Chondrogenic differentiation of clonal mouse embryonic cell line ATDC5 in vitro: Differentiationdependent gene expression of parathyroid hormone (PTH)/ PTH-related peptide receptor. J. Cell Biol. *133*, 457–468.
- Mortier, G.R., Cohn, D.H., Cormier-Daire, V., Hall, C., Krakow, D., Mundlos, S., Nishimura, G., Robertson, S., Sangiorgi, L., Savarirayan, R., et al. (2019). Nosology and classification of genetic skeletal disorders: 2019 revision. Am. J. Med. Genet. *179*, 2393–2419.
- **32.** Cheng, Y., and Patel, D.J. (2004). Crystallographic structure of the nuclease domain of 3'hExo, a DEDDh family member, bound to rAMP. J. Mol. Biol. *343*, 305–312.
- **33.** Thomas, M.F., L'Etoile, N.D., and Ansel, K.M. (2014). Eri1: A conserved enzyme at the crossroads of multiple RNA-processing pathways. Trends Genet. *30*, 298–307.
- 34. G Walleshauser III, J., Kessler, T., Morse, D., A Tannous, B., H L Chiu, N., and Chiu, H.L. (2012). A Simple Approach for Evaluating Total MicroRNA Extraction from Mouse Brain Tissues. J. Anal. Sci. Methods Instrum. 02, 5–12.
- **35.** Cox, T.C., Lidral, A.C., McCoy, J.C., Liu, H., Cox, L.L., Zhu, Y., Anderson, R.D., Moreno Uribe, L.M., Anand, D., Deng, M., et al. (2019). Mutations in GDF11 and the extracellular antagonist, Follistatin, as a likely cause of Mendelian forms of orofacial clefting in humans. Hum. Mutat. *40*, 1813–1825.
- 36. Gedeon, Á.K., Colley, A., Jamieson, R., Thompson, E.M., Rogers, J., Sillence, D., Tiller, G.E., Mulley, J.C., and Gécz, J. (1999). Identification of the gene (SEDL) causing X-linked spondyloepiphyseal dysplasia tarda. Nat. Genet. 22, 400–404.
- 37. Bamshad, M., Lin, R.C., Law, D.J., Watkins, W.C., Krakowiak, P.A., Moore, M.E., Franceschini, P., Lala, R., Holmes, L.B., Gebuhr, T.C., et al. (1997). Mutations in human TBX3 alter limb, apocrine and genital development in ulnar-mammary syndrome. Nat. Genet. 16, 311–315.
- **38.** Lv, D., Luo, Y., Yang, W., Cao, L., Wen, Y., Zhao, X., Sun, M., Lo, W.H.Y., and Zhang, X. (2009). A novel single-base deletion in ROR2 causes atypical brachydactyly type B1 with cutaneous syndactyly in a large Chinese family. J. Hum. Genet. *54*, 422–425.
- De Mori, R., Romani, M., D'Arrigo, S., Zaki, M.S., Lorefice, E., Tardivo, S., Biagini, T., Stanley, V., Musaev, D., Fluss, J., et al. (2017). Hypomorphic Recessive Variants in SUFU Impair the Sonic Hedgehog Pathway and Cause Joubert Syndrome with Craniofacial and Skeletal Defects. Am. J. Hum. Genet. *101*, 552–563.

- **40.** Bakrania, P., Efthymiou, M., Klein, J.C., Salt, A., Bunyan, D.J., Wyatt, A., Ponting, C.P., Martin, A., Williams, S., Lindley, V., et al. (2008). Mutations in BMP4 cause eye, brain, and digit developmental anomalies: overlap between the BMP4 and hedgehog signaling pathways. Am. J. Hum. Genet. *82*, 304–319.
- **41.** Elliott, A.M., and Evans, J.A. (2006). Genotype-phenotype correlations in mapped split hand foot malformation (SHFM) patients. Am. J. Med. Genet. A *140*, 1419–1427.
- 42. Chen, P., Tan, Z., Shek, H.T., Zhang, J.N., Zhou, Y., Yin, S., Dong, Z., Xu, J., Qiu, A., Dong, L., et al. (2022). Phenotypic Spectrum and Molecular Basis in a Chinese Cohort of Osteogenesis Imperfecta With Mutations in Type I Collagen. Front. Genet. 13, 816078.
- **43.** Alexiadis, A., Delidakis, C., and Kalantidis, K. (2017). *Snipper,* an *Eri1* homologue, affects histone mRNA abundance and is crucial for normal *Drosophila melanogaster* development. FEBS Lett. *591*, 2106–2120.
- **44.** Bohnsack, K.E., and Bohnsack, M.T. (2019). Uncovering the assembly pathway of human ribosomes and its emerging links to disease. EMBO J. *38*, e100278.
- **45.** Ridanpää, M., Van Eenennaam, H., Pelin, K., Chadwick, R., Johnson, C., Yuan, B., VanVenrooij, W., Pruijn, G., Salmela, R., Rockas, S., et al. (2001). Mutations in the RNA component of RNase MRP cause a pleiotropic human disease, cartilage-hair hypoplasia. Cell *104*, 195–203.
- 46. Vega, H., Waisfisz, Q., Gordillo, M., Sakai, N., Yanagihara, I., Yamada, M., Van Gosliga, D., Kayserili, H., Xu, C., Ozono, K., et al. (2005). Roberts syndrome is caused by mutations in ESCO2, a human homolog of yeast ECO1 that is essential for the establishment of sister chromatid cohesion. Nat. Genet. 37, 468–470.

- **47.** Boocock, G.R.B., Morrison, J.A., Popovic, M., Richards, N., Ellis, L., Durie, P.R., and Rommens, J.M. (2003). Mutations in SBDS are associated with Shwachman-Diamond syndrome. Nat. Genet. *33*, 97–101.
- 48. Ng, S.B., Buckingham, K.J., Lee, C., Bigham, A.W., Tabor, H.K., Dent, K.M., Huff, C.D., Shannon, P.T., Jabs, E.W., Nickerson, D.A., et al. (2010). Exome sequencing identifies the cause of a mendelian disorder. Nat. Genet. 42, 30–35.
- 49. Dixon, M.J. (1996). Treacher Collins syndrome. Hum. Mol. Genet. 5, 1391–1396.
- Farley-Barnes, K.I., Ogawa, L.M., and Baserga, S.J. (2019). Ribosomopathies: Old Concepts, New Controversies. Trends Genet. 35, 754–767.
- Carapito, R., Konantz, M., Paillard, C., Miao, Z., Pichot, A., Leduc, M.S., Yang, Y., Bergstrom, K.L., Mahoney, D.H., Shardy, D.L., et al. (2017). Mutations in signal recognition particle SRP54 cause syndromic neutropenia with Shwachman-Diamond-like features. J. Clin. Invest. *127*, 4090–4103.
- 52. Bellanné-Chantelot, C., Schmaltz-Panneau, B., Marty, C., Fenneteau, O., Callebaut, I., Clauin, S., Docet, A., Damaj, G.L., Leblanc, T., Pellier, I., et al. (2018). Mutations in the SRP54 gene cause severe congenital neutropenia as well as Shwachman-Diamond Like syndrome. Blood *132*, 1318–1331.
- **53.** Trainor, P.A., and Merrill, A.E. (2014). Ribosome biogenesis in skeletal development and the pathogenesis of skeletal disorders. Biochim. Biophys. Acta *1842*, 769–778.
- 54. Han, M., Chang, M., Kim, U.J., and Grunstein, M. (1987). Histone H2B repression causes cell-cycle-specific arrest in yeast: Effects on chromosomal segregation, replication, and transcription. Cell 48, 589–597.

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# Supplemental information

# Null and missense mutations of *ERI1* cause

## a recessive phenotypic dichotomy in humans

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## **Supplementary Note: Case Reports**

### **Individual 1A**

She was born at term and she had bilateral severe vesicoureteral reflux causing antenatally diagnosed hydronephrosis, which required bilateral reimplantation. She has asthma, partial conductive hearing loss, hypernasal speech, radial head dislocation, scoliosis and vertebral anomalies. She has chronic intermittent right hip pain, with decreased range of motion. She has joint stiffness especially in the morning, treated with ibuprofen. She has recurrent gingival infections. She has a past medical history of bronchitis, mastitis, and a Bartholin cyst. She has a normal intelligence. She enjoys cycling, talking walks and painting. She works as a babysitter and in a shop.

Examination: Height 112 cm (-8 SD). Weight 24 kg (-5 SD). Head circumference 50.6 cm (-3.5 SD). Hands 12 cm (-6 SD), feet 13 cm (-8 SD). Face is long and narrow. She has proptosis with slightly upslanting palpebral fissures and full eyebrows. Zygomatic hypoplasia. Prominent alveolar processes of the maxilla and mandibule. Facial features are somewhat coarse. The speech is hypernasal. The ears are low-set and small. Dolicocephaly. Limited elbow extension (150 degrees). Limited hip abduction. Joint hypermobility wrists and fingers. On the right, she has only one phalanx on the thumb; she has clinodactyly with ulnar deviation of the 2nd and 3rd fingers at the level of the PIP, she has a short 5th finger with clinodactyly (radial deviation). On the left hand, she has clinodactyly with ulnar deviation). She has pes planus with prominent heels. She has large 2nd toes, and syndactyly of the 2nd and 3rd toes on the right, and 4th and 5th toes on the left.

Radiology: Decreased tubulation of long bones with decreased medullary space. Hypoplastic femoral heads.Bilateral radioulnar subluxation. Dextroconvex lumbar scoliosis. Increased height of the vertebrae, with decreased anterior-posterior diameter. Narrow metacarpal bones. Bone mineral density at the lower limit of normal, no fractures.

There were no hematological anomalies.

# Individual 1B

Antenatally diagnosed hydronephrosis. Term delivery His birth weight was 4 pounds and his length was 14 <sup>3</sup>/<sub>4</sub> inches. He is known for bilateral severe vesicoureteral reflux causing severe hydronephrosis, impaired renal function and hypertension. Bilateral reimplantation of ureteras at age 3. He also has velopharyngeal insufficiency, a submucous cleft palate, mild conductive hearing

loss, asthma, 2 febrile seizures, congenital right hip dislocation which was repaired at 3 years of age. History of transient neutropenia, but otherwise no hematological anomalies. Bilaterally dislocatable patellae, operated at age 20. Some neck pain and ankle rigidity. Normal development and intelligence.

Physical examination. Height 128 cm (-7 SD), Weight 26 kg (-5 SD) Head circumference 53 cm (10th centile), Hands 15 cm (-3 SD), feet 17.5 cm (-5 SD). There is bitemporal narrowing. Proptosis with slightly upslanting palpebral fissures and full eyebrows. The chin is small but there are prominent alveolar processes of the maxilla and mandibule. The facial features are somewhat coarse. Zygomatic hypoplasia. The speech is hypernasal and the palate is high. The ears are lowset with prominauris. The chest is wide with mild pectus excavatum. The muscle mass is decreased. The elbows can't be extended beyond 150 degrees but there is wrist hypermobility. Beighton score of 5/9, with points for hypermobility at the level of the hands and back. There is extendable camptodactyly of the 2nd-5th fingers. Tapering fingers. Pes planus with prominent heels. There were no abnormalities on the neurological exam except for bilateraly brisk deep tendon reflexes. There is normal neck range of motion. Feet: left 4-5 cutaneous syndactyly.

Radiology: slight scoliosis with vertebral bodies of increased height, decreased anterio-posterior diameter, and irregular vertebral plates. The pelvis is dysplastic with right coxa vara and left coxa valga. The diaphyses are thin and the medullary space is narrow for multiple bones. There is hypoplasia of the olecranon process, the distal radial and ulnar epiphyses, the proximal radius and the carpal bones. There was bilateral dislocation of radio-ulnar joints and patellae. Narrow metacarpal bones with narrow medullary space.

# Individual 2

The Individual was a male neonate who was admitted to the neonatal ICU after birth due to respiratory distress. He was noted to have short limbs on prenatal ultrasound at 24 weeks. An amniocentesis was performed and testing for mutations in *FGFR3*, *COL1A1* and *COL1A2* did not identify rare variants.

At birth he was noted to be short, with a birth length of 40cm (-5.17SD). Additionally, he had several dysmorphic features: facial dysmorphic features included a prominent forehead, flat nasal bridge, micrognathia, large ears. He also had bilateral complete cutaneous syndactyly of fingers 3-5 and toes 4-5, 5th finger clinodactyly and a narrow thorax. Further testing included a chromosomal microarray and FLNB gene sequencing, which were both normal. No hematological abnormalities were noted. Due to pulmonary hypertension and right heart failure, he passed away at age 5 months. On autopsy, giant chondrocytes were seen in the epiphyseal plates of the long bones. No other systemic or organ malformations were noted.

# Individual 3

The boy was born to non-consanguineous Japanese parents at 39 weeks and 6 days of gestation. His birth length was 41.7 cm (-4.3 SD) and weight 2,000 g (-3.3 SD). He presented with tachypnea leading to a conventional respiratory support (nasal directional positive airway pressure). He showed oligosyndactyly of the right hand, proximally set left thumb, post-axial syndactyly of the feet, dislocation of the right elbow, and ulnar deviation of the left wrist. He was bottle-fed. His weight was 3,340 g at 3.5 months of age (< -3 SD). Respiratory support was weaned to nasal continuous positive airway pressure, and continued until hospital discharge at 4 months. Soon after then, he developed upper airway infection and subsequent respiratory failure, and came to an emergency room. Ultrasonography revealed pulmonary hypertension with severe tricuspid regurgitation. Despite ventilator support, he suddenly died the next day after hospital admission.

His X-rays showed platyspondyly and metaphyseal dysplasias of the long tubular bones. The left hand showed type A1 brachydactyly, while the right hand showed oligosyndactyly. Both feet showed post-axial syndactyly. The test of blood cells showed macrocytic anemia and congenital neutropenia.

# Individual 4

The male child was noted to have severe intrauterine growth retardation. He was born prematurely and had postnatal growth restriction with short stature. His past medical history included mild conductive hearing loss of left ear, delayed dentition with abnormal enamel and eruption, kidney hydronephrosis with grade 5 vesicoureteral reflux, small atrial septal defect, 2 small ventricular septal defects and a prominent aortic root. He had generalized hypotonia, delayed motor milestones, delayed speech. His failure to thrive was severe and he passed away at 26 months.

Examination showed dysmorphic facial features (trigonocephaly, frontal bossing, small jaw, sparse hair, small and low-set ear, posteriorly rotated, prominent pre-maxilla and a recessed chin retrognathia). He had puffy hands and feet, nonpitting edema, and undescended testes. Limb anomalies included dysplastic digits of both feet (rudimentary 2nd and 3rd toes and absent 4th and 5th toes on the left, right foot with 4th and 5th toe syndactyly)

Skeletal abnormalities including pectus carinatum deformity, skeletal dysplasia (osteopenia, wormian bones, severely delayed bone age).

There were no hematological anomalies.

## Individual 5

She is female born at term with a weight of 2.760 kg. There is no known parental consanguinity. Her past medical history includes bilateral inguinal hernias, a ventricular septal defect (perimembranous), and pulmonary vein stenosis. She has global developmental delay and autism. She was reported as F5 in a previous paper<sup>1</sup>, since she is homozygous for a *KYNU* variant (NM\_003937.3:c.616G>A, p.Glu206Lys) which affects a non-conserved amino acid but may perhaps still impact her clinical picture.

Examination at 8y and 9 mo: Weight 22 kg (8th centile) height 130.8 cm (46th centile), head 48.9 cm (just <2SD). Brachycephaly with a flat occiput. Bitemporal narrowing, small nose ears cupped and prominent. The right hand showed a transverse terminal limb reduction with a rudimentary 5th digit. Her left hand was less affected with hypoplastic and deep-set nails. Her right foot only had digits 1 and 5 which however displayed anonychia. The left foot showed an oligodactyly with digits 2 to 4 being absent. Digits 1 and 2 had syndactyly and anonychia.

There were no skeletal anomalies (normal spine, metaphyses and epiphyses) apart from the extremities.

She had a normal complete blood count at 11 months.

# Individual 6

He is a male born at 39 weeks (weight 2810 g (-1,6 SD), length 49 cm (-1,3 SD), head circumference 35 cm (-0,2 SD)). His parents are first cousins and he has a healthy younger brother. He has strabismus and mild intellectual disability, but is otherwise functioning well.

Examination at 13y 3m showed a height of 155 cm (25th centile) and height of 62 kg (85th centile) and a head circumference of 56 cm (75th centile). He has high arched eyebrows, a low-set forehead hairline and low-set ears. His hands showed a clinodactyly 5th finger bilaterally, brachydactyly, slightly dysplastic nails. Feet examination showed short wide feet, short, low set 4th left toe, very slight partial syndactyly 2-3.

Radiographies showed a normal skeleton apart from extremities. Hands: Brachymesophalangy 5, pseudoepiphyses, hypoplastic distal phalanges. Feet Short and thin left metatarsal 4, short left metatarsal 5

# Individual 7

The proband is the first child of consanguineous parents (first cousins) born at term by emergency Caesarean section due to fetal distress. Pregnancy itself was unremarkable. Birth weight was 2.6kg (0.4th centile). There were no concerns regarding the probands early development. The parents became concerned when she appeared not to be coping well at school. She is reported to have poor memory with limited concentration. She is sociable but mum has concerns that she can sometimes be inappropriately social towards strangers. She attends mainstream school with a statement of educational needs. At the age of 5 years 11 months (71 months), she underwent a Griffiths Mental Development assessment. Her developmental age at this time was 54.5 months for eye-hand coordination, 35 months for performance and 48 months for practical reasoning.

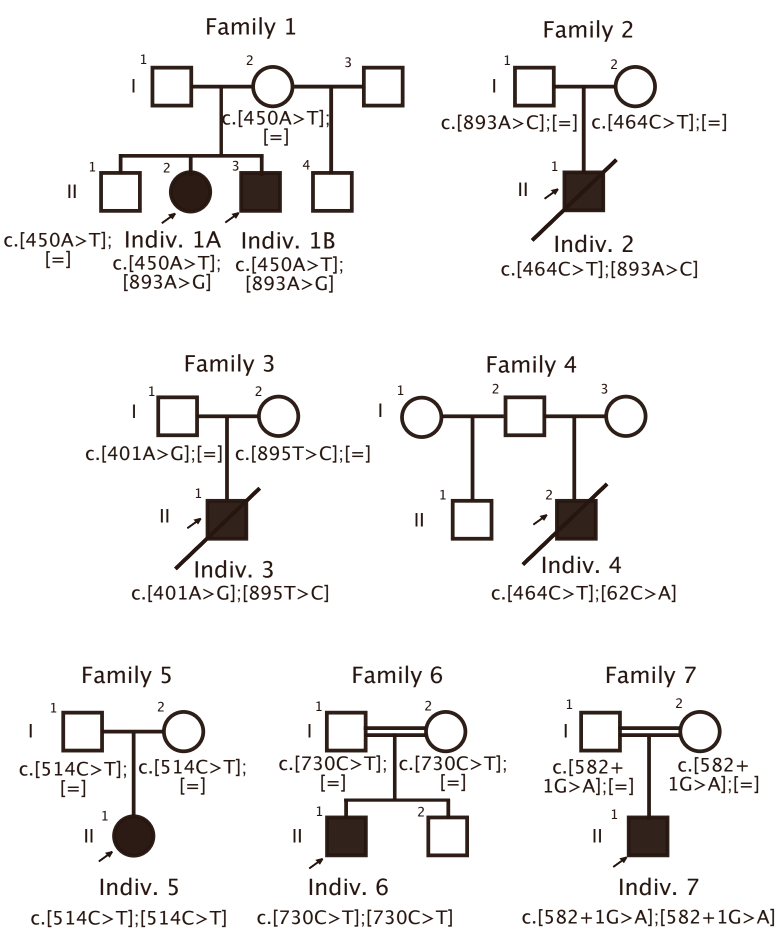
The proband had a urine infection when 5 months of age and she was subsequently found to have vesicoureteric reflux and hydronephrosis with scarring of her left kidney. She was commenced on prophylactic antibiotics.

On examination at 7 y, Weight 27.6 kg (50th centile) Height 130 cm (90th centile). She had prominent epicanthic folds and columella with upturned earlobes. She had shortening of the 5th metacarpals bilaterally and absence of the 5th toe on the right foot with overriding of the 3rd and 4th toes. The 4th toe on the left appeared absent and there was duplication of the 5th toe with syndactyly.

There were no radiological anomalies apart from those at the extremities. Abnormal appearance of the epiphyses of both distal radius and probably also distal ulna with overlapping proximal carpal bones. At age 8 years, bone age of 11 years on radiographs of hands and feet. Hands: Short right 1st metacarpal. Bilaterally short 5th metacarpals and mild shortening of 4th. Proximal phalanx right thumb is short. The right 5th finger distal phalanx is small with an abnormal epiphysis which is thickened.

Feet: Right: Missing the 5th ray. The fourth digit proximal phalanx and epiphysis are abnormal in shape. Overriding of the 3rd, 4th and 5th toes. Proximal 3rd metatarsal is fused to the 4th metatarsal. 4th metatarsal is widened. Left: There are five rays in the left foot. Proximal fusion of 4th and 5th metatarsals. Bilaterally: The 1st metatarsals are broadened and the shaft of the metatarsals and multiple epiphyses and phalanges have abnormal shapes in both feet.

Urine organic acids, urine amino acids, serum urea and electrolytes, full blood count, ferritin and thyroid function were all within normal limits.



**Figure S1.** Pedigrees of the families described. Variants are named according to NM\_153332.4.

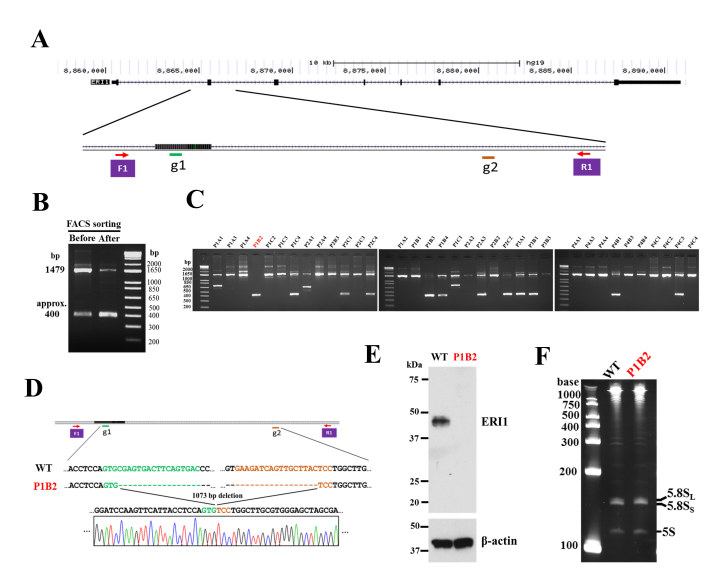
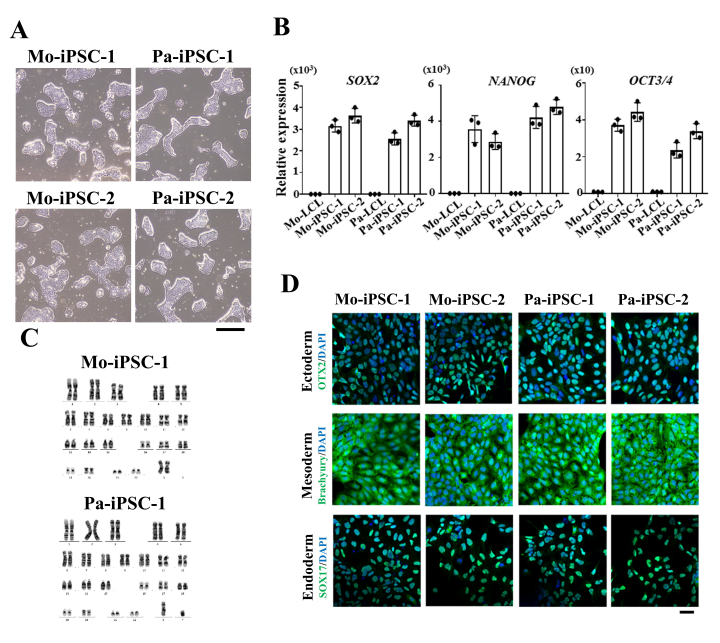
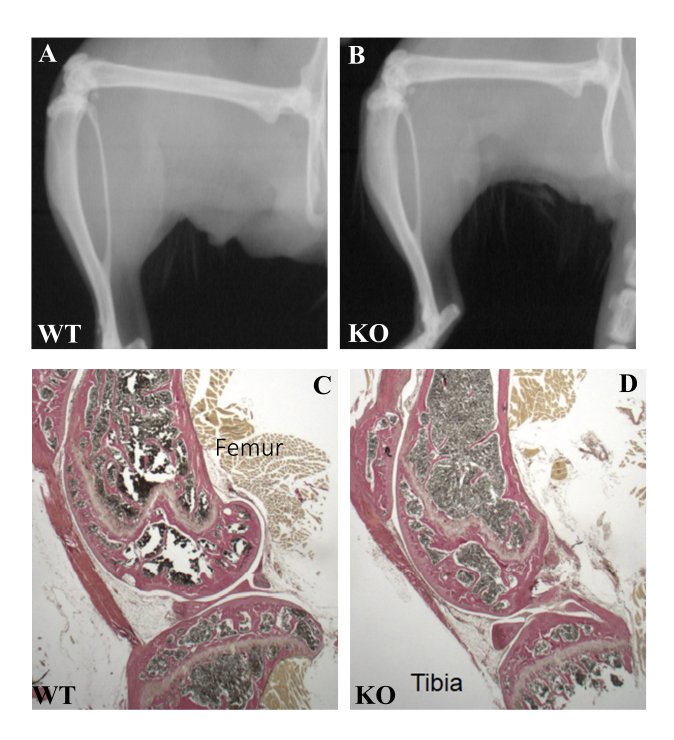


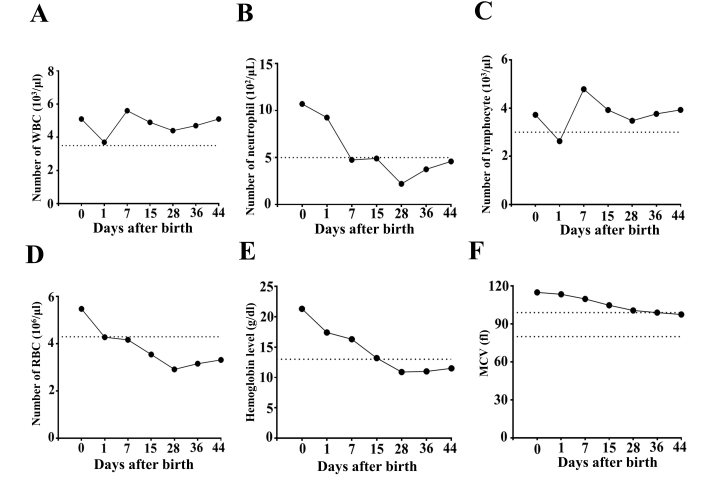
Figure S2. Generation of the *ER11* knock-out Hela cell line by CRISPR/Cas9mediated gene editing. A, Two guide RNAs (g1, 2) were designed to target genomic DNAs within exon 2 and intron 2 of *ER11*, respectively. Mixture of g1 and g2 were introduced into Hela cells with Cas9. **B**, The proportion of the cells with a genomic deletion generated by g1 and g2 was increased after FACS sorting. **C**, Monoclonal screening for *ER11* knock-out lines. The clone P1B2 without the PCR band corresponding to the wild type (marked in red) was selected. **D**, Sanger sequencing confirming a 1,073-bp genomic deletion in the P1B2 line. **E**, Western blot validation of the knock-out effect. **F**, Ethidium bromide-stained RNA.  $5.8S_L$ , long form of 5.8S rRNA;  $5.8S_S$ , short form of 5.8S rRNA.



**Figure S3. Characterization of induced pluripotent stem cell lines (iPSCs). A,** Representative bright-field views of iPSCs of the healthy mother (Mo-iPSC-1 and 2) and affected individual 3 (Pa-iPSC-1 and 2) generated from the lymphoblastoid cell lines (Mo- and Pa-LCLs). Scale bars, 500  $\mu$ m. **B,** RT-qPCR analysis showing the stem cell markers (*SOX2, NANOG* and *OCT3/4*) expressed in the iPSCs. n = 3 biologically independent samples. The data indicate mean  $\pm$  SD. **C,** Normal karyotypes in Mo- and Pa-iPSCs by chromosomal G-banding analysis. **D,** Evaluation of differentiation capacity into three germ layers of the iPSC lines by immunostaining for SOX17 (endoderm), Brachyury (mesoderm) and OTX2 (ectoderm). Scale bars; 50  $\mu$ m.



**Figure S4: Evaluation for the skeletal phenotype of hind limbs. A-B,** X-ray images of the wild type (WT) and *Eril* knockout (KO) mice at 14 weeks. Compared with WT mice, KO mice do not show significant morphological changes in the epimetaphyseal portions of the femurs and tibias. **C-D**, Van Gieson's staining for the epimetaphyseal portions of femur and tibia. KO mice do not show significant changes in the morphology and ossification of the epimetaphyses.



**Figure S5. Blood tests of Individual 3. A,** Number of white blood cells (WBCs). Normal range:  $3.5-8.5 [\times 10^3/\mu l]$ . The dotted line shows the lower limit of the normal value. **B**, Number of neutrophils. Normal range:  $[27-144] \times 10^2/\mu l$  from birth to 2 weeks; 10-50  $[\times 10^2/\mu l]$  since 2 weeks. The dotted line shows the diagnosis standard for severe congenital neutropenia. **C**, Number of lymphocytes. Normal range:  $3.0-9.5 [\times 10^3/\mu l]$ . The dotted line shows the lower limit of the normal value. **D**, Number of red blood cells (RBCs). Normal range:  $4.3-5.6 [\times 10^6/\mu l]$ . The dotted line shows the lower limit of the normal value. **E**, Hemoglobin level. Normal range: 13-17 [g/dl]. The dotted line shows the lower limit of normal value. **F**, Mean corpuscular volume (MCV). Normal range: 80-99 [fl]. The dotted lines show the lower and upper limits of the normal value.

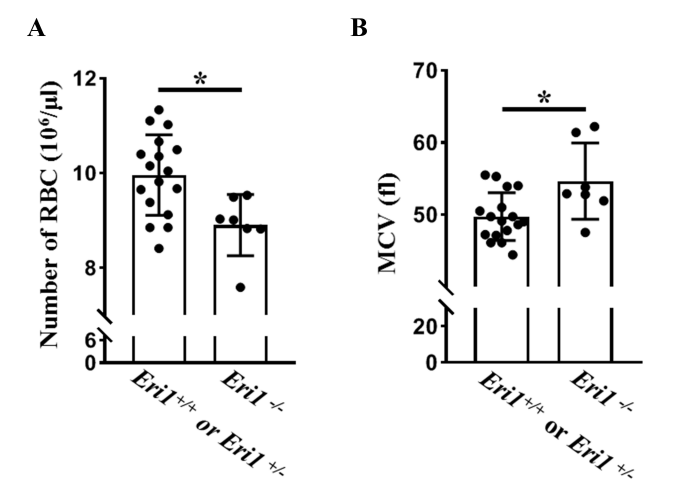


Figure S6. Blood tests of mice at 8 weeks. A, Number of red blood cells (RBCs).  $Eri1^{+/+}$  or  $Eri1^{+/-}$  mice, n=17;  $Eri1^{-/-}$  mice, n=7. \*P < 0.05. B, Mean corpuscular volume (MCV).  $Eri1^{+/+}$  or  $Eri1^{+/-}$  mice, n=17;  $Eri1^{-/-}$  mice, n=7. \*P < 0.05. The above quantitative data indicate mean  $\pm$  SD and statistical significance was assessed using two-sided t-test.

Primer name	Sequence (5'-3')					
Expression plasmid constru	ction					
Sal1 ERI1 CDS f	TCGGTCGACCATGGAGGATCCACAGAGTAAAGAGC					
Not1 ERI1 CDS r	TACGCGGCCGCGTTACTTTCTAAAATGTGGCATTTGTGGTG					
ERI1 D134G f	CCACTTGTGAAGAAGGAAACCCA					
ERI1 D134G r	CTTCAAAGcCAATAATACAAATGTAGTCATAATAACTGT					
ERI1 E150 f	TACATGAcATAATTGAATTTCCGGTTGTTTTACTG					
ERI1 E155 f	TACATGAAATAATTGAATTTCtGGTTGTTTTACTG					
ERI1 E150 P155 r	CAAACTCAGGTGGGTTTCCTTCT					
ERI1 S298 f	GATGgCTCTAAGAATATCGCCCGAATAG					
ERI1 S298 r	AAGACCACAGTGAGGCCGC					
ERI1 S299P f	GACcCTAAGAATATCGCCCGAATAGCAG					
ERI1 S299P r	ATCAAGACCACAGTGAGGCCG					
Sanger sequencing						
ERI1 EX1 f	TCAACGGAGAAAGGCGAG					
ERI1 EX1 r	TGAACACAGAATTAAGTTTAAAGCAA					
ERI1 EX3 f	ATGGTTTGAAGTACCAGTATATGTTGGAG					
ERI1 EX3 r	GATGTCAGAGGCCCTATCAGAGAAC					
ERI1 EX4 f	GCTGTGATGAATTCGTCGTG					
ERI1 EX4 r	CCAAGCAAGGTATTAAATTGTTCAC					
ERI1 EX6 f	TTTGTAGGAAGCAAGGTTAATTTTC					
ERI1 EX6 r	TTTGAACATAAGGAATTTGGAATC					
ERI1 EX7 f	CTGGTCTCATACTCCTGACCTCAAG					
ERI1 EX7 r	GCTGGTGACAAAGAATTCAGTATGCC					
Genome editing for Hela cel	lls					
ERI1 e2 g1 f	CACCgGTCACTGAAGTCACTCGCAC					
ERI1 e2 g1 r	AAACGTGCGAGTGACTTCAGTGACc					
ERI1 i2 g2 f	CACCgGAAGATCAGTTGCTTACTCC					
ERI1 i2 g2 r	AAACGGAGTAAGCAACTGATCTTCc					
PCR for ERI1 g1g2del f1	CATGTAGCATGAAGAGGCTTCAGAAG					
PCR for ERI1 g1g2del r1	TTCTCATGCTGTACCTGGCCAG					
RT-qPCR						
COL2A1_RT_F	GGTGGCTTCCATTTCAGCTA					

Supplementary table 1. Primers used in this study

COL2A1_RT_R	TACCGGTATGTTTCGTGCAG
OCT3/4_RT_F1	GACAGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGGG
OCT3/4_RT_R2	CTTCCCTCCAACCAGTTGCCCCAAAC
SOX2_RT_F	GGGAAATGGGAGGGGTGCAAAAGAGG
SOX2_RT_R	TTGCGTGAGTGTGGATGGGATTGGTG
NANOG_RT_F	CAGCCCCGATTCTTCCACCAGTCCC
NANOG_RT_R	CGGAAGATTCCCAGTCGGGTTCACC
ACTB_RT_F	AGAAAATCTGGCACCACACC
ACTB_RT_R	AGAGGCGTACAGGGATAGCA
HPRT_F	GCCTATAGACTATCAGTTCCCTTTGG
HPRT_R	TGCTGTGGTTTAAGAGAATTTTTTCA
HIST1H2AB_F	ACTCGGTCTTCTCGTGCAG
HIST1H2AB_R	GCTCCTCGTCATTGCGGAT
HIST2H2BE_F	CCGCAAAGAGAGCTACTCCA
HIST2H2BE_R	GTGGAGCGCTTGTTGTAGTG
HIST2H3C_F	CTACCAGAAGTCCACGGAGC
HIST2H3C_R	TGGATGGCGCACAGGTTC
HIST2H4A_F	GGCGGAAAAGGCTTAGGCAA
HIST2H4A_R	CCAGAGATCCGCTTAACGCC
HIST1H1A_F	TGGGCATTAAGAGCCTGGTAA
HIST1H1A_R	ACCCGTTGCCTTAGTTTTTGTA
TBP_F	TCTGGAATTGTACCGCAGCTT
TBP_R	GCTCCTGTGCACACCATTTT

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# References

1. Szot, J.O., Slavotinek, A., Chong, K., Brandau, O., Nezarati, M., Cueto-González, A.M., Patel, M.S., Devine, W.P., Rego, S., Acyinena, A.P., et al. (2021). New cases that expand the genotypic and phenotypic spectrum of Congenital NAD Deficiency Disorder. Hum Mutat *42*, 862-876. 10.1002/humu.24211.