Dear Editor,

Metaphyseal dysplasia without hypotrichosis (MDWH, OMIM #250460), a rare phenotype of cartilage-hair hypoplasia (CHH, OMIM #250250) - anauxetic dysplasia (AD, OMIM #607095) spectrum disorders, is described as an isolated skeletal dysplasia with no extra skeletal manifestations including hypoplastic hair and defective immunity [1]. Pathogenic variants in RNA component of the mitochondrial RNA-processing endoribonuclease (RMRP) gene cause CHH-AD spectrum disorders with autosomal recessive inheritance [2]. We report the first case in Korea of a girl with compound heterozygous RMRP variants, n.81G>A and n.100C>T, that were diagnosed as MDWH. The Catholic University of Korea St. Vincent's Hospital Institutional Review Board, Seoul, Korea, approved this study (number: VC-202ASEO0064).

A girl aged five years and 10 months was referred to the pediatric endocrine clinic at St. Vincent Hospital for short stature. She was the only child of non-consanguineous parents, born at 40 weeks and one day of gestation by spontaneous vaginal delivery, with a birth weight of 2.85 kg (10th–25th percentile), height of 45 cm (3th–5th percentile), and normal head circumference. The mid-parental height (maternal height, 166 cm; paternal height, 180 cm) was 166.5 cm. Her medical history did not reveal any immunodeficiency warning signs, and she had normal psychomotor development. On physical examination, her weight, measured using a CAS scale (CAS, Seoul, Korea), was 15.9 kg (-2.1 standard deviation score [SDS]) and her height, measured using a Harpenden Stadiometer (Holtain Ltd., Crymych, UK), was 96.4 cm (-4.2 SDS). The upper:lower segment ratio (1.3:1) was higher than the normal range for her age (Fig. 1). The patient's radiographic features revealed subtle cupping and widening at the metaphysis of both metacarpal bones, phalanges, and ulnae, as well as an expanded and irregular costochondral junction in her chest. The chest X-ray revealed a normal thymus. The peripheral blood examination results were as follows: leukocyte count, 5.79×10^9/L (neutrophils 49.8%, lymphocytes 39.9%); hemoglobin, 135 g/L (mean corpuscular volume [MCV] 85.4 fL); platelet count, 307×10^9/L; and erythrocyte sedimentation rate, 7 mm/h (normal range 0–20 mm/hr). The patient’s karyotype was 46,XX. The serum levels of calcium...
[median: 2.42 mmol/L (range: 2.1–2.59 mmol/L)], phosphorus [median: 1.48 mmol/L (range: 0.81–1.45 mmol/L)], and alkaline phosphatase (4.23 µkat/L (range: 0.5–2 µkat/L) showed no definite abnormalities. The thyroid and adrenal function tests were normal. The maximum serum growth hormone (GH) level following GH stimulation in the morning with levodopa 250 mg (Myung In Pharm., Seoul, Korea) and arginine 8 g (GREEN CROSS Well Being, Gyeonggi-do, Korea) was >10 µg/L. The patient was diagnosed as having idiopathic short stature, and GH treatment (0.3 mg/kg/week) was initiated. However, the patient’s Δheight SDS during the six months of GH therapy was 0.2; thus, we ceased GH treatment owing to poor growth response.

At six years and six months of age, after obtaining written informed consent, genomic DNA was extracted from the patient’s peripheral blood leukocytes, and a library was prepared using a custom designed panel (Celemics Inc., Seoul, Korea), which enriches a 616 kb region spanning 62 short-stature-related genes. Massive parallel sequencing was performed using the Illumina MiSeqDx platform (Illumina, San Diego, CA, USA). The targeted gene panel sequencing results revealed two variants in RMRP (NR_003051.3:n.81G>A and n.100C>T). Although neither parent consented to be tested for the RMRP variants, the two variants were confirmed as compound heterozygous based on the next generation sequencing results using the Illumina MiSeqDx platform. Therefore, the diagnosis was confirmed nine months after the patient's first visit. RMRP variants n.81G>A (previous nomenclature: c.99C>T) have both been reported in patients with CHH [3, 4] (Fig. 2).

The global incidence of MDWH is unknown; to date, only 20 MDWH patients have been confirmed by RMRP analysis [5-8]; over 90 RMRP variants have been identified in CHH-AD spectrum disorders [9]. Although the genetic and molecular background of the genotype-phenotype correlation in patients with RMRP variants is unclear, the position of the pathogenic variants in the transcript and the proposed effect on transcript folding and RNA/protein interaction result in the variable phenotypes observed in CHH-AD spectrum disorders [10]. As MDWH patients are deemed mildly affected, clinicians do not always consider potential extraskeletal complications. However, a study reported serious late onset bone marrow failure and cancer risks in MDWH patients [6]. Clinicians should consider mild-phenotype MDWH, even in patients exhibiting only short stature.

**ACKNOWLEDGEMENTS**

None.

**AUTHOR CONTRIBUTIONS**

Conceptualization: Cho WK; Analysis: Ki CS, Hyun H, Yoo IH, Cho WK, and Namgoong H; and Writing - review and editing: Cho WK, Lee JH, and Oh JH.
Fig. 2. Integrated genome browser snapshot of the RNA component of mitochondrial RNA-processing endoribonuclease (RMRP) gene variants. The orientation of RMRP is in the reverse direction, so that the n.100C>T variant is on the left and the n.81G>A variant is on the right. The distance between both variants is 19 bp, meaning that many of the variants exist on the same sequence reads, thus confirming the compound heterozygosity of the two variants. The red dotted boxes indicate sequence reads with only the n.100C>T variant, while the blue dotted boxes indicate sequence reads with only the n.81G>A variant.
CONFLICTS OF INTEREST

No potential conflicts of interest relevant to this article were reported.

RESEARCH FUNDING

None declared.

ORCID

Ho Namgoong https://orcid.org/0000-0001-7295-2455
Chang-Seok Ki https://orcid.org/0000-0001-7679-8731
Hyesun Hyun https://orcid.org/0000-0001-8525-1471
Il Han Yoo https://orcid.org/0000-0002-7438-6925
Won Kyoung Cho https://orcid.org/0000-0003-0918-0565
Jung Hyun Lee https://orcid.org/0000-0002-1849-733X
Jin-Hee Oh https://orcid.org/0000-0002-2893-0563

REFERENCES

1. Verloes A, Pierard GE, Le Merrer M, Maroteaux P. Recessive metaphyseal dysplasia without hypotrichosis. A syndrome clinically distinct from McKusick cartilage-hair hypoplasia. J Med Genet 1990;27:693-6.
2. Horn D, Rupprecht E, Kunze J, Spranger J. Anauxetic dysplasia, a spondyloepiphyseal dysplasia with extreme dwarfism. J Med Genet 2001; 38:262-5.
3. Hermanns P, Tran A, Munivez E, Carter S, Zabel B, Lee B, et al. RMRP mutations in cartilage-hair hypoplasia. Am J Med Genet A 2006;140: 2121-30.
4. Reicherter K, Veeramani AI, Jagadess S. Cartilage-hair hypoplasia caused by novel compound heterozygous RMRP mutations. Indian Pediatr 2011; 48:559-61.
5. Bonafé L, Schmitt K, Eich G, Giedion A, Superti-Furga A. RMRP gene sequence analysis confirms a cartilage-hair hypoplasia variant with only skeletal manifestations and reveals a high density of single-nucleotide polymorphisms. Clin Genet 2002;61:146-51.
6. Vakkilainen S, Costantini A, Taskinen M, Wartiovaara-Kauto U, Mäkitie O. ‘Metaphyseal dysplasia without hypotrichosis’ can present with late-onset extraskeletal manifestations. J Med Genet 2020;57:18-22.
7. Nakashima E, Mabuchi A, Kashimada K, Onishi T, Zhang J, Ohashi H, et al. RMRP mutations in Japanese patients with cartilage-hair hypoplasia. Am J Med Genet A 2003;123A:253-6.
8. Bonafé L, Dermitzakis ET, Unger S, Greenberg CR, Campos-Xavier BA, Zankl A, et al. Evolutionary comparison provides evidence for pathogenicity of RMRP mutations. PLoS Genet 2005;1:e47.
9. Thiel CT and Rauch A. The molecular basis of the cartilage-hair hypoplasia-anauxetic dysplasia spectrum. Best Pract Res Clin Endocrinol Metab 2011;25:131-42.
10. Mäkitie O and Vakkilainen S. Cartilage-hair hypoplasia - anauxetic dysplasia spectrum disorders. In: Adam MP, Ardinger HH, et al., eds. GeneReviews [Internet]. Seattle, WA: University of Washington, Seattle. 2012.