**POLR1C variants dysregulate splicing and cause hypomyelinating leukodystrophy**

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**Abstract**

**Objective**

To further clarify the molecular pathogenesis of RNA polymerase III (Pol III)-related leukodystrophy caused by biallelic POLR1C variants at a cellular level and potential effects on its downstream genes.

**Methods**

Exome analysis and molecular functional studies using cell expression and long-read sequencing analyses were performed on 1 family with hypomyelinating leukodystrophy showing no clinical and MRI findings characteristic of Pol III–related leukodystrophy other than hypomyelination.

**Results**

Biallelic novel POLR1C alterations, c.167T>A, p.M56K and c.595A>T, p.I199F, were identified as causal variants. Functional analyses showed that these variants not only resulted in altered protein subcellular localization and decreased protein expression but also caused abnormal inclusion of introns in 85% of the POLR1C transcripts in patient cells. Unexpectedly, allelic segregation analysis in each carrier parent revealed that each heterozygous variant also caused the inclusion of introns on both mutant and wild-type alleles. These findings suggest that the abnormal splicing is not direct consequences of the variants, but rather reflect the downstream effect of the variants in dysregulating splicing of POLR1C, and potentially other target genes.

**Conclusions**

The lack of characteristic clinical findings in this family confirmed the broad clinical spectrum of Pol III–related leukodystrophy. Molecular studies suggested that dysregulation of splicing is the potential downstream pathomechanism for POLR1C variants.

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RNA polymerase III (Pol III)-related leukodystrophy is characterized by hypomyelination in the CNS with various additional manifestations such as hypogonadotropic hypogonadism, hypodontia, cerebellar ataxia, and atrophy of the corpus callosum. After the proposal of multiple clinical entities, the discovery of pathogenic variants in genes encoding 2 major subunits of Pol III, POLR3A and POLR3B, in the majority of patients led to the concept of Pol III–related leukodystrophy being emerged. Recently, variants in yet another gene coding for Pol III complex, POLR1C, were identified in patients who were negative for, but showed clinical features similar to those with, POLR3A and POLR3B variants.

Here, we report a patient with novel POLR1C pathogenic variants, who showed clinical and imaging features compatible with hypomyelinating leukodystrophy without additional features characteristic of Pol III–related leukodystrophy. We also propose a potential molecular mechanism of POLR1C variants involving dysregulation of splicing.

Methods

This study was approved by the Institutional Review Board of the National Center of Neurology and Psychiatry. Genomic DNA and total RNA were extracted from the peripheral blood of the patient and parents. For DNA diagnostic testing, we performed quantitative PCR for the screening of PLP1 duplication, followed by exome sequencing for Mendelian disease panel (TruSight One, Illumina), as we previously performed according to the manufacturer’s protocol. POLR1C complementary DNAs were obtained by reverse transcriptase (RT)-PCR, which were cloned into an expression vector, pCMV3.1 (Invitrogen) with FLAG-tag at the N-terminus, for subsequent fluorescent immunostaining. For long-read next-generation sequencing, barcoded RT-PCR products (control, father, mother, and patient) were sequenced on a single MinION R9.4 flow cell (Nanopore).

Data availability

Any data not published within the article will be shared by request from any qualified investigator.

Case report

The patient was a Japanese boy without a family history of neuromuscular diseases and had normal neurodevelopment during infancy. At age 2 years, he developed action tremor of his fingers, had difficulty in writing, and showed early signs of motor dyspraxia. At age 3 years, he developed amblyopia secondary to hypermetropia and astigmatism. Myopia was not noted. At age 3 years and 10 months, he presented with action tremors in fingers, but there were no other neurologic abnormalities. He showed a developmental quotient of 105 (Enjoji analytical developmental test for infants and toddlers). Subsequently, he became neurodevelopmental abilities stagnated and regressed in his daily activities and nystagmus became apparent. On his visit at age 5 years and 9 months, he had a short stature with 101.4 cm (–2.1 SD) tall without apparent microcephaly, facial abnormalities, or ambiguous genitalia. No delay or abnormal order in dentation was noted. He exhibited lateral nystagmus, action tremors, and slurred speech. The finger-to-nose, pronosupination, and tandem walk tests showed mild dysmetria and ataxia. Deep tendon reflexes of the lower limbs were increased. He exhibited a staggering wide-based gait and was unable to stand on 1 leg for more than 2 seconds. Both parents were intellectually and physically normal with no neurologic findings.

We performed several tests at age 5–6 years. Laboratory tests revealed prepubertal patterns of pituitary gonadotropins and testosterone. The Wechsler Intelligence Scale for Children, fourth edition, showed regression with a full-scale intelligence quotient of 70. Peripheral nerve conduction velocities and auditory brainstem responses were normal. EEG was normal. MRI showed diffuse T2 hyper- and T1 iso-intensities in the white matter, indicating hypomyelination (figure 1, A and B). T1 and T2 shortening in the optic radiation, the ventrolateral thalamus, and the dentate nucleus was noted, as typically observed in Pol III–related leukodystrophy (figure 1, A–C). Cerebellar atrophy or thinning of the corpus callosum was not evident (figure 1, C and D). MRIs of the parents were not available.

Results

After PLP1 duplication was excluded, the panel exome sequencing identified 2 novel heterozygous missense variants in exon 3 and exon 6 of the POLR1C gene (NM_203290.3: c.167T>A, p.M56K and c.595A>T, p.I199F, respectively; figure 1E). Parental segregation analysis confirmed compound heterozygosity. In silico prediction analyses revealed both variants to be pathogenic at different levels (figure 1F). RT-PCR using the patient’s sample showed increased proportion of splicing variants with a combination of full intron 3 and/or half/full intron 4 inclusions, all of which are presumably nonfunctioning variants with premature termination codons on sequence validation (figure 2, A and B). The patient’s major transcript was the variant including both intron 3 and intron 4. Three representative variants expressed in HeLa cells showed that p.M56K alone did not change the protein stability, but the nuclear localization was modestly diminished (figure 2, C–G). p.M56K with intron 3 and intron 4 inclusion significantly decreased the protein level. Meanwhile, p.I199F
caused cytosolic punctation and reduced protein expression (figure 2, C–H). The punctation did not overlap with lysosomal marker Lamp1, an autophagosome marker LC3, or with the proteosome marker, ubiquitin (data not shown). To our surprise, both parents also showed increased proportion of the intron 3/4 inclusion variant (figure 2A), which prompted us to use long-read next-generation sequencing to obtain deep reads of all variants with allelic segregation. Mapping patient POLR1C transcripts on genome demonstrated that they were biallelic, and more than 85% of correctly mapped transcripts were intron-containing variants (figure 3A). Both parents also showed apparently increased proportion of intron-containing variants (64% in the father and 52% in the mother). These findings suggested 2 possible mechanisms: (1) both c.167T>A and c.595A>T variants directly affected the splicing to properly remove intron 3/4, or (2) splicing abnormality resulted from impaired function of Pol III target genes that play roles in the maintenance of splicing. To delineate these 2 possibilities, the parental reads of each allele were remapped to determine whether each variant affected splicing in cis or trans. Surprisingly, the proportion of intron-containing variants was equivalent between wild-type and mutant alleles in both parents, indicating that both variants affect splicing in trans (figure 3B). Because it is unlikely that the variant in one allele directly affects the splicing of the other, this trans effect is probably driven by the latter hypothesis.

Discussion

In this study, we reported 1 family with novel POLR1C compound heterozygous variants. Clinically, our patient so far had no characteristic features of Pol III–related leukodystrophy, such as dental abnormality and hypogonadism, and no cerebellar atrophy and thinning of the corpus callosum. Although these features are common in patients with POLR3A or POLR3B variants, thereby serving as diagnostic key features, findings other than hypomyelination are not necessarily
The molecular mechanisms underlying POLR1C variants causing hypomyelination remain unknown. We showed that the POLR1C variant on each allele does not only affect the subcellular localization and/or amount of the protein but also affects the splicing that removes the intron 3/4. Allelic present in all cases.\textsuperscript{10} Previously reported 32 cases with POLR1C variants presented with at least one of these features.\textsuperscript{6,11,12} Thus, the present case also suggested that lacking characteristic features of Pol III–related leukodystrophy does not exclude the presence of POLR1C variants.

Agarose gel electrophoresis images of RT-PCR products using total RNA from blood cells, amplifying (A) the entire length and (B) exon 2–exon 5 of POLR1C cDNA. (A) Although a normal control (NL) shows a single 1-Kb band (white arrowhead), the patient (PT) showed longer abnormal bands (red arrowheads) along with a fainter normal band. Both the father (FA) and the mother (MO) also showed abnormal bands with different intensities. (B) In addition to the expected 340-bp normal band (white arrowhead), 3 additional bands were observed in both the normal and patient samples (red arrowheads). Band isolation and sequencing confirmed that the strongest amplicon in the patient (842 bp band) included both intron 3 and intron 4 (int3 + int4). The others contain either the entire or the first half of intron 4 (int4 or int4-half, respectively). (C) A scheme of each variant transcript cloned into an expression vector, pCDNA3.1. Arrowheads indicate each variant. M2 includes intron 3 and intron 4 (int3 + int4) flanking exon 3. (D) Western blot of POLR1C variants transiently expressed in HeLa cells. Upper panel: POLR1C; middle panel: enhanced green fluorescent protein (EGFP) (cotransfected as an inner control for normalization of transfection efficiency); lower panel: actin (loading control). Cells were harvested after 24 hours of transfection. An anti-FLAG antibody was used to visualize exogenous POLR1C. The sizes of the M2 band appear to be the same as WT, suggesting that these introns are partially spliced out before translation. Truncated protein was not observed, presumably due to the removal by nonsense-mediated messenger RNA decay before translation. (E) Quantification of the POLR1C protein level. Experiments were performed in triplicates. POLR1C was normalized to EGFP. The y-axis indicates relative value to the average of wild type. (F) fluorescent immunostaining of HeLa cells transiently expressing wild-type and mutant POLR1C. Subcellular localization of exogenous protein was determined using anti-FLAG antibody. Bar indicates 20 μm. Wild-type POLR1C showed strong nuclear expression (WT). The M56K mutant showed reduced nuclear staining (M1). The I199F mutant showed prominent cytosolic punctations (yellow arrowheads; M3). DAPI nuclear staining shows blue signals. Images were obtained using Keyence BZ-X710 fluorescence microscope (Keyence, Japan). (G) Quantification of nuclear localization. Using imaging software (Keyence), the ratio of signal intensities of the nucleus and cytosol was measured (n = 30 cells per group). (H) The proportion of cells with cytosolic punctations was calculated in more than 300 cells (10 fields [30–40 cells per field] at 200× magnification). Error bars indicated standard errors. *p < 0.05, **p < 0.01, ***p < 0.001. One-way analysis of variance.
Figure 3 Depth of coverage of aligned reads of POLR1C cDNA

(A) Top: schematic representation of POLR1C consisting of 9 exons (black rectangles). We examined transcript variant 1 (NM_203290.3). Bottom: the bar graph shows the sequence depth at each position, and the number shows the average depth of exons and introns. Red numbers highlight aberrant transcripts. In addition to intron 3 and intron 4 inclusion, skipping of exon 7 was evident in a small proportion of transcripts. From the top: control (blue), father (green), mother (light blue), and patient (yellow). The proportion of aberrant splicing variants was obtained by dividing the highest read count of intron 4 by the highest read count of all exons (e.g., 77879/91464 in the patient). (B) Alignment of each allelic reads. The c.167T and c.167A allele reads were selected from sequence reads of the father, and the c.595A and c.595T allele reads were selected from those of the mother. Ten thousand reads of each allele were aligned. There was no obvious difference in the proportion of variants with intron inclusions between the 2 alleles in each parent.
segregation studies using a long-read sequencing technology in the patient revealed a large proportion of abnormal splicing variants. Moreover, the analyses in the parental samples showed 2 unexpected findings. First, in addition to the exon 3 variant, the exon 6 variant also affected splicing even in a heterozygous status, indicating that haploinsufficiency of \textit{POLR1C} caused molecular deficits despite the autosomal recessive mode of inheritance. Second, each allelic variant caused the inclusion of introns on both alleles. This most likely resulted from altered transcription of Pol III target genes that play a role in the regulation of splicing of \textit{POLR1C}. As one such candidate, we examined the expression of U6 snRNA, which plays a central role in splicing-some,\textsuperscript{13} but was not altered in either the patient nor his parents (data not shown). Although the exact effectors downstream of the \textit{POLR1C} variants remain unknown, our findings provide a potential molecular mechanism for \textit{POLR1C} variants that affect the activity of Pol III and transcription of its target genes, which may lead to dysregulation of splicing of genes including \textit{POLR1C}.

In conclusion, we reported 1 family with hypomyelinating leukodystrophy caused by novel \textit{POLR1C} variants. Both pathogenic variants resulted in changes in subcellular localization and reduction in protein levels, as well as inclusion of introns, which presumably resulted in loss of function. Allelic segregation analyses of full-length transcripts in both patient and parents revealed that the aberrant splicing variants are not direct consequences of the coding variants, but rather reflect the downstream effect of the variants in dysregulating splicing of \textit{POLR1C}, and potentially other target genes.

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\textbf{Disclosure}

The authors report no disclosures. Go to Neurology.org/NG for full disclosures.

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**References**


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