Metachromatic leukodystrophy: Characterization of two (p.Leu433Val, p.Gly449Arg) arylsulfatase A mutations

YANGYANG WANG, XIANG CHEN, CHAN LIU, SHAMIN WU, QINGFENG XIE, QUAN HU, SHAN CHEN and YIWEI LIU

Department of Rehabilitation, The Second Affiliated Hospital and Yuying Children's Hospital of Wenzhou Medical University, Wenzhou, Zhejiang 325000, P.R. China

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Abstract. Metachromatic leukodystrophy disorder (MLD) is an autosomal recessive lysosomal storage disease. The disease is primarily caused by a deficiency in the enzyme arylsulfatase A (ASA), which is encoded by the ARSA gene. A total of 254 mutations have been reported in different populations. The present study aimed to detect causative gene mutations in an atypical case presenting with attention deficit hyperactivity disorder through whole-exome sequencing. Of note, the patient’s mother is from a consanguineous family. Compound heterozygous variants (c.1297C>G) + (c.1345G>A) [(p.Leu433Val) + (p.Gly449Arg)] were identified in exon 8 in the ARSA gene of the pediatric patient. The two missense mutations identified have not been previously reported, to the best of our knowledge. Furthermore, an in silico analysis and multiple phylogenetic tree analyses of ARSA homologs were performed to predict the effects of the two novel mutations. Serial changes were observed in the patient with MLD at follow-up visits over 6 years. However, brain MRI images demonstrated no notable progression and the number of ASA enzymes was stable. Also, the results of neurodevelopmental assessment showed that the patient was diagnose with ADHD. These data may offer a potential explanation of the genotype-phenotype correlation in MLD and enhance the spectrum of mutations associated with the condition.

Introduction

Metachromatic leukodystrophy (MLD; Mendelian Inheritance in Man #250100) is an autosomal recessive inherited leukoencephalopathy characterized by demyelination of the central and peripheral nervous systems (1). These pathological changes are caused by failure of sulfatides and other glycolipid-containing sulfuric acids to be desulfurized and their subsequent deposition in the lysosomes of systemic tissues. Desulfurization requires the combined efforts of the enzyme arylsulfatase A (ASA; Enzyme Commission no. 3.1.6.8), encoded by the ARSA gene, a lysosomal hydrolase, and sphingolipid activator protein B, encoded by the PSAP gene. A deficiency in any of these components results in abnormal metabolism of sulfatides, which triggers progressive degenerative metabolic encephalopathy (2,3). The worldwide prevalence of MLD is estimated to be between 1 in 40,000 and 1 in 170,000 (4).

MLD is divided into three clinical subtypes according to the age at onset of the disease: Late infantile (<4 years), juvenile (4-14 years) and adult (>14 years) (5). Clinical manifestations of MLD are varied and lack specificity (6). Three types of MLD are differentiated by the presence or absence of neurological symptoms and the pattern of disease progression. The severity of MLD appears to be negatively correlated with the age at onset (7). Infantile MLD is the most common form, accounting for ~60% of all diagnosed cases. The first clinical symptoms of infantile MLD are mainly the deterioration of the motor system, including spastic tetraparesis, frequent falls and walking on toes. As the disease progresses, a patient may develop flaccid paralysis and lose the ability to stand, exhibit speech deficits and a decrease in psychological functions, develop optic atrophy, suffer generalized or partial seizures and develop peripheral neuropathy, causing death within 2-4 years from initial diagnosis (8-10). The symptoms of juvenile MLD include poor intellectual capabilities, emotional problems, language disorders and a gradual regression in motor function. Patients with juvenile MLD die within 10-15 years after the first symptoms appear, with only few cases reaching their 20th birthday (11,12). The adult form of the disease presents...
with neuromuscular or behavioural problems and progresses slowly (12).

A genotype-phenotype correlation has been reported in MLD (13-15). A deficiency in ASA is caused by mutations in the ARSA gene. The ARSA gene (GenBank accession no. NG_009260) is located on chromosome 22q13.33. This small gene (~3.5 kb) has eight exons and encodes 509 amino acid precursors (GenBank accession nos. NM_000487.5 and NP_000478.3). Studies have indicated that in all populations, the ARSA gene has three alleles (15-18). The first allele is named ARSA-MLD (a pair of pathogenic alleles). Polten et al (15) reported that the ARSA-MLD gene has two different alleles. Mutations in one allele, designated allele I (or O), result in non-functional activity of ASA. Mutations in the other allele, designated allele A (or R), maintain the residual activity of ASA. Patients who are homozygous for allele I or allele A mainly present with the late infantile form or the adult form of MLD, respectively. Patients who are heterozygous for the two alleles (I/A) usually present with the juvenile form (6). The second allele is a pair of ASA pseudodeficiency (PD) alleles (ARSA-PD). Whether the mutation is homozygous (ARSA-PD/ARSA-PD) or heterozygous (ARSA-PD/ARSA-MLD) (17), the level of ASA may be 5-15% of the reference value in certain patients with the mutant ARSA-PD gene, while the structure of ASA is moderately altered (19). In patients with a low level of ASA, residual enzyme activity is sufficient for normal physiological functions, which is known as PD. In European and American populations, the most common ARSA-PD allele has two variants (16). One is c.1049A>G (N350S), which prevents the ASA protein from entering lysosomes. The other is c.966A>G, a mutation in the 3′ untranslated region (poly A tail), which causes a significant reduction in ASA protein production. However, Asian populations have only one allele (c.1049A>G) of ARSA-PD. These Asian patients account for 20-30% of all patients with MLD (20). Furthermore, homozygosity for the mutant gene has little effect on the activity of ASA. The third allele is (ARSA-MLD; ARSA-PD) (18). It is reported that ARSA-MLD and ARSA-PD, known as the ARSA-MLD-PD allele, are located on the same chromosome. Based on the aforementioned genetic heterogeneity of MLD, it is required to identify this multiple allele in the ARSA gene to determine the genotype-phenotype correlation. To date, 254 mutations have been detected in the ARSA gene (21).

In the present study, whole-exome sequencing (WES) was performed in the family of a case of MLD to identify causative mutations and to offer a potential explanation for the genotype-phenotype correlation in MLD.

Subjects and methods

Subjects and clinical evaluation. The patient of the present study was recruited at Yuying Children's Hospital of Wenzhou Medical University (Wenzhou, China). At the first medical examination of the male patient at the age of 2 years 8 months due to a motor vehicle collision without any obvious indication of resulting neurological damage, a brain magnetic resonance imaging (MRI) scan was performed to ensure no underlying brain injury. The imaging unexpectedly revealed signs of MLD. Further enquiries regarding the proband's family history unveiled that the proband's grandparents (maternal parents) had a consanguineous marriage. Given the special family history, cognitive assessments were performed using the Gesell Development Scales test (22). It is the most common test to assess neurobehavioural development in infants (<3 years of age) through five factors (gross motor function, fine motor function, speech, adaptive behaviour and interpersonal behaviour). The results of Gesell Development Scales test was determined as development quotient (DQ) (22). The patient was evaluated by determining the DQ at the age of 2 years 8 months. Pediatric subjects scoring DQ<75, 75<DQ<85 and DQ≥85 exhibit low, marginal and normal development, respectively (23). In addition, several complementary inspections were performed to confirm the diagnosis, including ASA activity in leukocytes, electromyography (EMG) and gas chromatography-mass spectrometry (GC-MS) (24) at the proband's first visit. GC-MS represents an unbiased and open approach that allows the detection of unexpected changes in metabolite levels.

During 6 years of follow-up, the patient was subjected to continuous physical assessments. The Wechsler Intelligence Scale-third Edition (WAIS-III) test for neurodevelopmental assessment (25,26) was performed when the patient turned 4 years old. At our institution, the WAIS-III test is commonly used to evaluate the neurodevelopmental status of individuals aged ≥4 years (26). The results of the WAIS-III test are expressed as an intelligence quotient (IQ), and the IQ is scored according to the UK WAIS-III manual scoring criteria (25). An IQ of 90 is the boundary between normal and low intelligence. The WAIS-III comprises 14 subtests (vocabulary, similarities, information, comprehension, block design, matrix reasoning, picture completion, picture arrangement, coding, symbol search, digit span, letter-number sequencing, arithmetics and object assembly) (27). The Ayres Sensory Integration (ASI) test (28-30) was performed at the age of 7 years 11 months (July, 2017) to assess behavioural problems. Furthermore, neuroradiological studies were performed by brain MRI scans as the patient aged. At the most recent visit, the proband and the proband's parents were subjected to genetic testing.

WES. Genomic DNA was isolated from peripheral blood leukocytes of the affected proband and the proband's parents by Kangso Medical Inspection. Whole-exome capture using the SureSelect Human All Exon kit (Agilent Technologies, Inc.) and high-throughput sequencing were performed in-house as previously described (7,8). The reads were aligned for single-nucleotide variant (SNV) calling and subsequent analysis for prioritization of candidate genes (9). ANNOVAR (version 20180118) (31) was used to annotate the detected variations. ANNOVAR can utilize annotation databases conforming to Generic Feature Format version 3 (GFF3). The gene mutations we verified based on the low frequency of the detected variants provided in relevant gene databases [mainly the 1000 Genomes Project (https://www.ncbi.nlm.nih.gov/variation/tools/1000genomes/) and dbSNP (https://www.ncbi.nlm.nih.gov/SNP/)]. First, DNA samples were compared with the hg 19 reference (one version of normal human genome from NCBI database). If the detected mutations did not match the template file, a database was consulted (mainly the 1000 Genomes Project). Subsequently, according to the frequency
of base mutations given in the database, it was determined whether the mutation site is pathogenic. Furthermore, several studies suggest that in numerous cases, variations in the chromosome copy number (CNV), rearrangement and structure may be connected with disease (32). Therefore, the CNV was also determined by using CODEX (33) and XHMM software (34). CODEX and XHMM present a novel normalization and CNV calling method. CODEX was used to remove biases and artifacts in WES data, and produce accurate CNV calls. The XHMM software was designed to recover information on CNVs from targeted exome sequence data.

In silico analysis. Deleterious missense SNVs were predicted by the following web-based tools: i) Sorting Intolerant From Tolerant (SIFT; sift.bii.a-star.edu.sg), where an SNP with a SIFT score <0.05 predicts a negative effect on the encoded amino acid, ii) polyphen-2 (genetics.bwh.harvard.edu/pph2/), where an SNP with a score between 0.85 and 1.0 predicts a damaging effect on the encoded amino acid, while an SNP with a score between 0.0 and 0.15 is predicted to be benign and an SNP with a score between 0.15 and 1.0 predicts a possibly damaging function, and iii) Mutation Taster (www.mutation-taster.org), where a score close to 1 indicates a high ‘security’ of the prediction of the given variant to be disease-causing.

In addition, multiple phylogenetic tree analyses of ARSA homologs were performed using MEGA7 software (35).

Results

Clinical manifestation. A Chinese pedigree from the city of Wenzhou with an index patient (MLD01) was recruited for the study (Fig. 1). The diagnosis of MLD was made pre-symptomatically at the age of 2 years 8 months based on the typical presentation of brain MRI. At first, the patient was asymptomatic with generally normal development. Later, the proband exhibited mild behavioural problems and was diagnosed with attention deficit hyperactivity disorder (ADHD) at the age of 7 years 11 months based on the ASI examination. He had a normal gait with no signs of paraparesis, spasticity or neuropathy, but was socially immature based on his interactions with the surroundings and individuals, as well as the ASI test. He was in a relatively stable pre-onset stage at the time of writing. The result of the EMG was normal. To date, the proband has developed no other neurological symptoms and lives an ordinary life, except for suffering from ADHD and struggling to keep up with his classmates in a regular school.

ARSA activity determination. No biochemical abnormalities were detected in the proband. ARSA activity in leukocytes of the proband was 150 nmol/17 h/mg protein at the first visit (at the age of 3 years in 2012) and 142 nmol/17 h/mg protein at the most recent visit (at the age of 8 years in 2017). Notably, the activity of ASA in 2017 was at the lower end of the normal range (normal range: 134.1-325.1 nmol/17 h/mg protein).

Neurodevelopmental assessment. At the first visit at the age of 2 years 8 months in March 2012, testing on the Gesell Development Scale revealed a borderline DQ level (75.1). At 4 years of age (in 2013), WAIS-III testing (36) suggested that the patient had mild mental retardation (IQ, 81). Neurocognitive evaluation in 2014 indicated a slight abnormality in full-scale IQ with slow processing, low average working memory and below-average executive skills. On the last follow-up (in 2017), the IQ had reached an almost stable level at 80. In addition, ASI testing suggested that the patient presented with inattention and bad temper, and the patient was subsequently diagnosed with ADHD. Given the mild symptoms, it was suggested that the patient could focus on home education. The parents could have greater patience and provide sufficient attention to the child. In addition, attention exercises, including handwriting, were encouraged.

Neuroradiological studies. The images of the first MRI scan (Fig. 2Aa-c) in March 2012 exhibited a lightly enlarged lateral ventricle and sporadic tigroid pattern of white matter in the centrum semiovale and basal ganglia. These results...
are characteristic of MLD. On the last MRI scan (Fig. 2Ba-c) in 2017, white-matter hyperintensities were effectively unchanged. Over the 6-year follow-up period, brain MRI scans revealed no evidence of progressive demyelination, indicating that the patient is in a stable condition.

Mutation detection. The mutation frequency may be expressed as that in Asian populations (ASN) or in East Asians (EAS). The mutation frequency of ‘c.1297C>G’ in the ARSA gene of the proband was ASN=0.0035 and EAS=0.0020, respectively, and the mutation frequency of ‘c.1345G>A’ in the ARSA gene of the proband was identical (ASN=0.0035 and EAS=0.0020). Therefore, after variant filtering, the study focused on the compound heterozygous variants (c.1297C>G) + (c.1345G>A) in exon 8 in the ARSA gene of the patient, which result in p.Leu433Val and p.Gly449Arg amino acid substitutions, respectively. Sanger sequencing was performed to confirm the above results and to investigate the possible familial segregation of the mutations. Compound heterozygous mutations detected in the proband’s parents indicated that each of the proband’s parents was a heterozygous carrier of either of the two mutations (Fig. 3), who coincidentally passed the mutation on to the proband individually. Furthermore, no mutations with an autosomal-dominant pattern of inheritance were identified.

In addition, no suspicious CNVs were identified in the WES data through the aforementioned computational methods.

In silico analysis. To further characterize the missense mutations identified in the family, different in silico tools
were used to assess the influences that the novel c.1297C>G and c.1345G>A missense mutations have on ARSA protein function. Pathogenicity predictions by the Mutation Taster programme suggested that the c.1297C>G mutation is deleterious (Table I). Pathogenicity predictions by the SIFT and PolyPhen-2 programme suggested that the c.1297C>G mutation is tolerated and probably damaging, respectively. The accuracy of Mutation Taster is reportedly the highest of the three utilized tools (37).

Furthermore, it is thought that highly conserved amino acid sequences have a functional value and are significant for protein structure, which indicates that they have an important role in determining the conformation of different domains of a protein (38). Multiple sequence alignment of ARSA orthologues revealed that the amino acid L433 is highly conserved across mammalian species, from human to rat, and the amino acid G449 is less conserved. The evolutionary conservation of the amino acid L433 indicates its significance in the structure of the ASA protein (Fig. 4).

**Discussion**

The present study reported on a Chinese pediatric patient diagnosed with MLD manifesting as ADHD as the major clinical symptom. Of note, two missense mutations were identified in the ARSA gene of the patient. The compound heterozygous variants in the proband [c.1297C>G] + [c.1345G>A] were inherited from the proband's mother and father, respectively. Of note, no de novo variants were detected in the proband. According to software prediction and comparison of homologous proteins, C.1297 was indicated to be the major mutation contributing to the onset of the disease. By contrast, the patient's mother, who possesses the same mutation, is normal. From the phenotype, the proband had a tigroid pattern on every brain MRI and was diagnosed with mild behavioural problems with MLD, whereas the proband's parents had no symptoms. Notably, the subtype of adult MLD (>14 years) is characterized by neuromuscular or behavioural problems. However, the reported case showed behavioural problems at the age of 7 years. Therefore, it is likely that this presentation was due to the compound heterozygous variants [c.1297C>G] + [c.1345G>A].

Biochemical analysis indicated that the activity of ASA was normal in the proband, which indicates that the mutations in the two genes did not affect the enzyme level. Although test for ASA activity in 2017 was at the lower end of the normal range, ASA activity remained normal in patient's body. That is, there is no qualitative change in ASA. Combined with the results of a previous study, it may be hypothesized that c.1297C>G and c.1345G>A probably belong to mutations in the ARSA-PD allele. It is reported that the compound heterozygous mutation in ARSA-PD/ARSA-MLD is not associated with progressive neurological disease (17).

Neurologically, all brain MRI scans for the proband display pathological changes of demyelination, which is characteristic of MLD. Furthermore, the patient's MRI images exhibit little progression during the 6-year follow-up period. It may be hypothesized that this may be associated with the normal enzyme levels in the proband.

Concerning the phenotype, the patient's condition cannot be regarded as severe when compared to other cases of MLD.
Given that the disease is usually classified according to the age of onset, it is difficult to classify this case into a proper subtype of MLD. The patient presented with ADHD-like symptoms but had no neurological signs at the age of 7 years 11 months. The MRI images exhibited demyelination of the cerebrum at the age of 2 years 8 months. If this case was to be classified according to the age of onset, the patient should be diagnosed with juvenile MLD, according to the clinical symptoms, the patient, who displayed behavioural problems, should be diagnosed with the adult form. Therefore, subjects with suspicion of MLD should be comprehensively evaluated. The clinical manifestation of MLD is highly variable. If clinicians diagnosed MLD only based on the general indicators of the three subtypes, numerous patients would be misdiagnosed or the diagnosis would be missed. Subjects with suspected MLD, particularly in a consanguineous family, may be diagnosed through genetic testing. In addition, the phenotype is the result of interactions between genes and the environment. A tigroid pattern of white matter at the level of the centrum semiovale and basal ganglia hyperintensity indicated in MRI highly suggests a diagnosis of MLD, especially when considering the consanguinity relationship between the child’s grandparents. The clinical presentation of the proband of the present study is atypical, which may be associated with his living environment. Of note, he possessed a large number of good educational resources and his parents provided a nurturing environment and sufficient attention.

However, the present study only included one family. It may be worthwhile to perform a cohort study to further examine genotype-phenotype associations in MLD. In addition, experimental animal studies with the same mutations are required to investigate the biochemical characterization of MLD (39). By enlarging the number of samples for mutation analysis, it is possible to screen those subjects at risk or with symptoms to diagnose early. The authors of the current study may be able to summarise some common features by increasing the size of the cohort in future studies. As physician can recognize people at high risk of MLD, only individual gene test need to be performed rather than WES. This is helpful for detecting ARSA gene mutations quickly and reliably, and reducing the cost of genetic testing.

In conclusion, the compound heterozygous variants (c.1297C>G) + (c.1345G>A) were described in a Chinese pediatric patient with MLD. The patient, having normal activity of ARSA, presented with ADHD-like symptoms, no neurological signs but had no neurological signs at the age of 7 years 11 months. The MRI images exhibited demyelination of the cerebrum at the age of 2 years 8 months. If this case was to be classified according to the age of onset, the patient should be diagnosed with juvenile MLD, according to the clinical symptoms, the patient, who displayed behavioural problems, should be diagnosed with the adult form. Therefore, subjects with suspicion of MLD should be comprehensively evaluated. The clinical manifestation of MLD is highly variable. If clinicians diagnosed MLD only based on the general indicators of the three subtypes, numerous patients would be misdiagnosed or the diagnosis would be missed. Subjects with suspected MLD, particularly in a consanguineous family, may be diagnosed through genetic testing. In addition, the phenotype is the result of interactions between genes and the environment. A tigroid pattern of white matter at the level of the centrum semiovale and basal ganglia hyperintensity indicated in MRI highly suggests a diagnosis of MLD, especially when considering the consanguinity relationship between the child’s grandparents. The clinical presentation of the proband of the present study is atypical, which may be associated with his living environment. Of note, he possessed a large number of good educational resources and his parents provided a nurturing environment and sufficient attention.

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