INTRODUCTION

Solute carrier family 9 member A6 gene (SLC9A6, OMIM 300231) is located at Xq26.3 and encodes the alkali cation (Na⁺,K⁺)/proton (H⁺) exchanger NHE6 isoform protein, which plays a critical role in the regulation of pH lumen in early recycling endosomes and the process of neuronal arborization and synapse development.1–3 The alteration of SLC9A6 causes a phenotype mimicking Angelman Syndrome, now referred to as Christianson syndrome (CS,OMIM 300243).4 To date, at least 35 different SLC9A6 pathogenic variants have been identified.

CS is an X-linked neurodevelopmental and neurological disorder, characterized in males by critical symptoms that include intellectual disability, epilepsy, hyperkinesia, truncal ataxia, and postnatal microcephaly.

Background: Variants in the endosomal solute carrier family 9 member A6 (SLC9A6)/(Na⁺,K⁺)/H⁺ exchanger 6 (NHE6) gene have been linked to epilepsy, speech loss, truncal ataxia, hyperkinesia, and postnatal microcephaly.

Methods: In the present study, we evaluated genetic alterations in a 3-year-old Chinese boy displayed features of epilepsy, psychomotor retardation, microcephaly, low body weight, difficulty in feeding, excessive movement, attention loss, ataxia, and cerebellar atrophy and his healthy family using WES method. The identified variant was further confirmed by Sanger sequencing method. Finally, minigene assays were used to verify whether the novel SLC9A6 intronic variant influenced the normal splicing of mRNA.

Results: We identified a novel hemizygous splicing variant [NM_001042537.1: c.1463-1G>A] in SLC9A6 by trio-based exome sequencing. The minigene expression in vitro confirmed the splicing variant altered a consensus splice acceptor site of SLC9A6 intron 11, resulting in skipping over exon 12.

Conclusions: Our finding extends the catalog of pathogenic intronic variants affecting SLC9A6 pre-mRNA splicing and provides a basis for the genetic diagnosis of CS.

KEYWORDS
chinese boy, christianson syndrome, novel splicing variant, SLC9A6
microcephaly. Furthermore, these core phenotypic features can also be accompanied by secondary symptoms, such as autistic behaviors, poor weight gain, feeding difficulties, motor regression, sleep disturbances, squint, high pain threshold, hypotonia, gastroesophageal reflux, and different degrees of cerebellar atrophy (CA). Contrast to male, female variant carriers show milder phenotypes with variable penetrance. Affected CS patients show a remarkable neurological symptom may be due to high-level expression of mutant \( SLC9A6 \) gene in the central nervous system.

To date, few Chinese CS patients have been reported. Here, we report a Chinese boy presenting with Christianson syndrome due to novel splicing variant NM_001042537.1: c.1463-1G>A in the \( SLC9A6 \) gene. The variant is not mentioned in ClinVar, dbSNP, 1000genomes, NCBI, ExAC, or HGMD database. We performed minigene splicing assay to validate the functional effect of variant in \( SLC9A6 \) gene to determine whether the hemizygous site is a pathogenic site for the affected boy.

2 | MATERIALS AND METHODS

2.1 | Patient and ethics

The present study was approved by the Ethics Committee of Northwest women’s and children’s hospital (Xi’an, China). Written informed consent was obtained from the parents.

2.2 | Samples and DNA extraction

Genomic DNA was extracted from 2–3 ml EDTA peripheral blood sample using BloodGen Midi Kit (CWBio), according to the manufacturer’s protocol.

2.3 | Variant screening

Library enrichment for whole exome sequencing was conducted using SeqCap EZ Exome v3 kit (Roche NimbleGen).Then, the enriched samples were sequenced by using an Illumina Hiseq 2500 (Illumina). The variants which is presented at least 20% of all reads were identified as quality variants. Thus, all quality variants were evaluated by bioinformatic tools including Polyphen2, PROVEAN, and MaxEntScan. The pathogenicity of all quality variants was accessed by The American College of Medical Genetics and Genomics (ACMG) Standards and Guidelines. The pathogenic variant was validated by Sanger sequencing.

2.4 | Transcription analysis

\( SLC9A6 \) fragments containing intron11, exon12, and intron12 were amplified from DNA of patient and control by nested PCR. The first PCR was performed using genomic DNA as a template, with 37527-SLC9A6-F and 39956-SLC9A6-R as primers; the second PCR was performed using products from the first PCR as a template, with 37833-SLC9A6-F and 39651-SLC9A6-R as primers; the third PCR was performed using PCR products from the second round as a template, with pcMINI-SLC9A6-Kpn1-F and pcMINI-SLC9A6-BamHI-R as primers. The primer sequences were listed as Table 1.

Restriction enzymes Kpn I and BamH I were used to digest amplified products, and the digestion products were ligated to pcMINI plasmid to obtain the pcMINI-SLC9A6-wt and pcMINI-SLC9A6-mut plasmid (Figure 4A). The recombinant plasmids were then transfected into human embryonic kidney cells (HEK-293T) and human cervical cancer cells (HeLa) by lipofectamine 2000 (Invitrogen). After 48 h, cells were harvested and RNA was extracted by RNAiso Plus (TAKARA). cDNA was synthesized using PrimeScript RT Master Mix (TAKARA). Finally, the cDNA was identified using 1.8% agarose gel electrophoresis and verified through sequencing.

3 | RESULT

3.1 | Clinical case report

The boy was born in full-term normal delivery without anoxia and asphyxia history. His birth weight, length, and occipitofrontal circumference (OFC) were reported to be within normal range. There were no family histories of mental illness, genetic illness, epilepsy, and febrile seizures. His six-month-old sister was in good health. His parents were both Han Chinese.

He started a seizure at the age of 12 months, characterized by general tonic-clonic seizure, and has been treated with topiramate (TPM) firstly. TPM was stopped using without permission from doctor after half a month. However, 3 months later, he had frequent general tonic-clonic seizures in clusters for 6 months, despite antiepileptic treatment by sodium valprorate (VPA). General tonic-clonic seizures were suddenly completely controlled by additional use of levoethialacetam (LEV) and repetitive transcranial magnetic stimulation (rTMS), with a seizure-free interval between 22 months old and 34 months old. During the absence of seizures, rTMS has been added only a few times and then discontinued, LEV was replaced with lamotrigine (LTG) because of his hyperkinesis and bad appetite after using 6 months, and LTG was subsequently replaced with

| Primers | Sequence (5’-3’) |
|---------|-----------------|
| 37527-SLC9A6-F | ggcctagcaacaatatgctg |
| 37833-SLC9A6-F | cctcgccttttcgcggctg |
| 39651-SLC9A6-R | ttttatctgcaccttggtt |
| 39956-SLC9A6-R | gttttgaggttggtag |
| pcMINI-SLC9A6-Kpn1-F | ggtagtgattacacatggc |
| pcMINI-SLC9A6-BamHI-R | tagtgagttccatgctcatatat |
zonisamide (ZNS) because of his anaphylaxis after using one month. But nocturnal seizures were found at age of 34 months, characterized by consciousness loss and upward gaze, without other abnormal movements. Oxcarbazepine (OXZ) was added to the antiepileptic treatment in external hospital. However, general tonic-clonic seizures in clusters were appeared again at the age of 39 months, subsequently followed by myoclonic and atonic seizure in clusters. Just then, he was admitted to our hospital.

He was initially diagnosed as psychomotor retardation at the age of 9 months, due to a visible delaying of developmental milestones, including unstable sitting posture and undeveloped language. The first electroencephalograph (EEG) and magnetic resonance imaging (MRI) indicated nothing abnormal. And he received rehabilitation training regularly and achieved independent ambulation at the age of 2 years and 6 months finally, but his gait was insecure and ataxic. He had no autistic behaviors so far, but had hyperactive and distracted behaviors. He spoke little, occasionally, some unconscious reduplicated words. He had poor weight gain because of having poor appetite and frequent gastrointestinal discomfort. He hardly ever drooled.

On physical examination, his weight was 11 kg (<1st centile), belonged to severe underweight, his OFC 45 cm (<1st centile), belonged to microcephaly. His mandibular tension was high and with facial obesity, often closed teeth, restricted opening mouth. The neurological examination was unremarkable except the ataxic gait.

During hospitalization, most of the laboratory test results were normal, which included ammonia, homocysteine, inorganic phosphorus, serum calcium, 25-(OH)D, ceruloplasmin, thyroid function, erythrocyte sedimentation rate (ESR), urine organic acids analysis, and karyotype. Thus, only serum free carnitine concentration slightly reduced (7.68 μmol/L, normal range: 8.5–50 μmol/L).

Furthermore, EEG and MRI were rechecked. Awake EEG revealed that on the basis of diffuse irregular medium to high amplitude 4–6 Hz θ rhythm, middle to very high amplitude spikes, multiple spikes, spikes and slow waves complex, and rhythms were exploded frequently and asynchronously. The leads of frontal, occipital, and temporal were prominent. The right hemisphere was more obvious than the left. During sleep, epileptiform discharge was significantly increased than awake (see Figure 1). The brain MRI revealed hypoplasia of inferior parts of cerebellar vermis, and enlargement of inferior orifice of the fourth ventricle, and abnormal signals in the left frontal cortex. The other intracranial structures including brainstem, basal ganglia, and supratentorial brain structures were within normal limits (Figure 2).

Since he came out of our hospital, there has been a seizure-free interval of 5 months. During this period, his ambulation and appetite were a little better than before, but his language development and weight gain still changed little. He still had no autistic behavior, but he was easily irritated. He was more likely to suffer from upper respiratory infections than normal children, sometimes even had fever, but there was no seizure. General tonic-clonic, myoclonic, and atonic seizures were no longer appeared. But sleep seizures occurred occasionally again after 5 months, characterized by consciousness loss and upward gaze, without other abnormal movements, and relieved after about 3 minutes. The dose of LEV was adjusted from 30 to 60 mg/kg every day. After each dose adjustment, the frequency of seizures would be significantly reduced, but sleep seizures were not controlled completely.

3.2 | The whole exome sequencing reveals a novel SLC9A6 splicing variant

Whole exome sequencing was performed on the proband and his parents and sister, and variant filtering was performed as described in the methods. Annotated whole exome sequencing data were examined for variants in genes for relevance for the epilepsy and developmental delay phenotypes. Finally, one splicing variant was chosen as potential causal variant. The splicing variant NM_001042537.1: c.1463-1G>A was not previously reported in variant databases including ClinVar, dbSNP, 1000genomes, NCBI, ExAC, and HGMD database. Sanger sequencing confirmed the hemizygous variant of the SLC9A6 in the proband II-1, which was a de novo variant not founded in his parents and sister (Figure 3).
3.3 The intron c.1463-1G > A variant causes a skipping transcription of exon 12

In vitro transcription analysis was performed to validate how the variant affect splicing products. Figure 4B showed that mutant plasmid encoded a shorter transcriptional products compared to the WT plasmid. The results of Sanger sequencing demonstrated the intron mutant lead to a complete skipping of exon 12 (Figure 4C, Figure 4D).

4 DISCUSSION

According to our literature review and this study, at least 35 different SLC9A6 pathogenic variants have been identified, which included 16 exonic variants and 10 intronic splices site variants (Table 2). The exonic mutational spectrum showed that exon 12 was a high incidence region of pathogenic variants, accounting for 43.8% of all exonic pathogenic variants. SLC9A6 is a twelve-span transmembrane protein and exon 12 encodes the last transmembrane motif. The mutational spectrum indicated a vital role of exon 12 in the function of SLC9A6. The de novo splice-site variant [NM_001042537.1: c.1463-1G > A] in SLC9A6 was identified as a novel pathogenic variant in this report. This variant is closest variant to C-terminal of the known pathogenic intron variant, and it causes the skipping transcription of exon 12. Given the importance of exon 12, the pathogenicity of this variant is predicated.

This patient presented drug refractory epilepsy, global psychomotor development delay, microcephaly, underweight, feeding difficulties, hyperkinesis, attention-deficit, ataxic gait and mild CA, no mouth opening, and no drooling and had a de novo splice-site variant in SLC9A6 on the X chromosome which caused a skipping transcription of exon 12. Consequently, the patient was diagnosed as CS definitely.

The clinical presentations of this patient were almost consistent with the characteristics of CS, except for no mouth opening, no drooling, and no autistic behavior; many previous literatures also have described the above symptoms as secondary symptom. The detailed mechanism might be related to different variants of SLC9A6 gene leading to different phenotypes of CS.
FIGURE 3  A. Family’s pedigree and variant. Pedigree of the family. B. Sanger sequencing revealed a splice-site variant in SLC9A6 gene.

FIGURE 4  Splicing analysis using a minigene assay. (A) Construction of the pcMINI-SLC9A6-wt and pcMINI-SLC9A6-mut plasmid. (B) Reverse-transcription polymerase chain reaction (RT-PCR) products were separated by electrophoresis of the pcMINI-SLC26A4-wt/mut vector in HeLa and 293T cells. The longer band represented the wild-type transcript (WT) with a length of 543 bp, the shorter band represented the mutated transcript (MT) with a length of 249 bp. (C) Schematic diagram of minigene construction and schematic diagram of Sanger sequencing of RT-PCR products. (D) Sanger sequencing of the RT-PCR products, the wild-type transcript consisted of exon A, 12, and B, in contrast, the mutated transcript consisted of exon A and B, without exon 12.
| Exon | Alteration region | Alteration type | Variants (default transcript NM_001042537) | Protein alteration | Clinical feature | References |
|------|------------------|-----------------|------------------------------------------|-------------------|------------------|------------|
| 1    | Nonsense variant  | NM_006359: c.190G>T | p. Glu64* | Mental retardation, epilepsy, ataxia, drooling | Pescosolido et al., 2014 |
| 1    | Missense variant  | c.316A>G         | p. Met106Val | Mild mental retardation, severe behavioral disturbances | Ibarluzea et al. 2020 |
| 2    | Frame shift variant | c.441delG | p. Ser147fs | Microcephaly, dysphasia, epilepsy, ataxia, strabismus, drooling, no cerebellar atrophy | Takahashi et al. 2011 |
| 3    | Frame shift variant | NM_006359: c.477_481del | p. Ile160Leufs*5 | Severe mental retardation, generalized tonic seizure, microcephaly, cerebellar atrophy, ataxia, hypotonia | Ikeda et al. 2019 |
| 3    | Frame shift variant | c.582_595del   | p. Tyr194fs | Focal seizures (impaired awareness), no cerebellar atrophy | Liu et al. 2018 |
| 4    | Frame shift variant | NM_006359: c.512_513delAT | p. His171fs | Mental retardation, microcephaly, epilepsy, ophthalmoplegia, squint, motor regression, ataxia | Christianson et al., 1999 |
| 4    | Frame shift variant | c.540_547dupAGAAGTAT | p. Phe183fs*1 | Mental retardation, epilepsy, microcephaly, ataxia, drooling, autism, hypotonia, no cerebellar atrophy | Pescosolido et al., 2014 |
| 5    | Frame shift variant | NM_006359: c.608del | p. His203Leufs*10 | Developmental delay, seizure, movement disorder, dystonia | Trump et al. 2016 |
| 6    | Non-frame shift variant | NM_006359: c.764_769delAAAGTG | p. Glu255_Ser256del | Developmental delay, epilepsy,ataxia, microcephaly, verbal language absent, easily provoked laughter | Gilfillan et al., 2008 |
| 6    | Frame shift variant | c.838_839delGinsG | p. Leu280Alafs*17 | Severe mental retardation, microcephaly, generalized seizure, autism | Fung et al. 2017 |
| 7    | Non-frame shift variant | c.1012_1020del | p. Trp338_Thr340del | Severe mental retardation, dysphasia, autism, ataxia, motor regression, unilateral weakness, no cerebellar atrophy | Garbern et al., 2010 |
| 7    | Nonsense variant | c.916C>T | p. Gln306* | Mental retardation, microcephaly, cerebellar atrophy, epilepsy, ophthalmoplegia, squint, drooling, flexed arms, motor regression, ataxia, dystonia | Mignot et al., 2013 |
| 9    | Nonsense variant | c.1219C>T | p. Gln407* | Mental retardation, epilepsy, microcephaly, ataxia, drooling, autism, hypotonia, cerebellar atrophy | Schroer et al., 2010 |
| 9    | Frame shift variant | NM_006359: c.1222_1226del | p. His408Asns*2 | Early infantile epileptic encephalopathy, dystonia | Trump et al. 2016 |
| 10   | Splice-site variant and missense variant | c.1148G>A | p. Gly383Asp | Mental retardation, epilepsy, microcephaly, ataxia, drooling, autism, no cerebellar atrophy | Pescosolido et al., 2014 |
| 11   | Frame shift variant | c.1414dupA | p. Arg472fs*4 | Mental retardation, epilepsy, microcephaly, ataxia, drooling, autism, hypotonia, cerebellar atrophy | Pescosolido et al., 2014 |
| 12   | Nonsense variant | c.1498C>T | p. Arg468* | Developmental delay, epilepsy,ataxia, microcephaly, verbal language absent, easily provoked laughter, no cerebellar atrophy | Gilfillan et al., 2008 |
| 12   | Frame shift variant | NM_006359: c.1464_1465insT | p. Thr489Tyrfs*23 | Severe mental retardation, microcephaly, seizure, scoliosis | Riess et al., 2012 |

(Continues)
| Alteration region | Alteration type | Variants (default transcript NM_001042537) | Protein alteration | Clinical feature | References |
|-------------------|----------------|-------------------------------------------|-------------------|-----------------|------------|
| Exon 12           | Frame shift variant | c.1505_1509dupCTGCC | p. Thr504Leufs*8 | Mental retardation, suspicion of epilepsy, microcephaly | Yalcintepe et al. 2021 |
| Exon 12           | Nonsense variant | c.1568G>A | P. Trp523* | Mental retardation, epilepsy, microcephaly, ataxia, drooling, autism, hypotonia, no cerebellar atrophy | Pescosolido et al., 2014 |
| Exon 12           | Nonsense variant | c.1569G>A | P. Trp523* | Mental retardation, epilepsy, electrical status epilepticus in sleep, autism | Mathieu et al., 2018 |
| Exon 13           | Nonsense variant | c.1639G>T | p. Glu547* | Mental retardation, epilepsy, microcephaly, ataxia, happy behavior | Schuurs-Hoeijmakers et al., 2013 |
| Exon 14           | Nonsense variant | c.1710G>A | P. Trp570* | Mental retardation, epilepsy, microcephaly, ataxia, drooling, autism, hypotonia, | Pescosolido et al., 2014 |
| Exon 15           | Missense variant | NM_006359: c.1831G>A | p. Glu611Lys | Global developmental delay, cerebellar atrophy, focal and generalized tonic-clonic seizures, hypotonia, large head | Padmanabha et al. 2017 |
| Intron 2           | Splice-site variant | c.526-1,G>A | - | Mental retardation, epilepsy, microcephaly, ataxia, drooling, autism, hypotonia, cerebellar atrophy | Pescosolido et al., 2014 |
| Intron 2           | Splice-site variant | c.526-9_526-5del | Skipping of exon 3 (mRNA validation) | Mild Mental retardation, Dysphasia, No autistic behavior | Masurel-Paulet A et al. 2016 |
| Intron 3           | Splice-site variant | NM_006359: c.507+1delGTAA | p.V144_R169del | Developmental delay, epilepsy,ataxia,microcephaly, verbal language absent, easily provoked laughter, no cerebellar atrophy | Gillilan et al., 2008 |
| Intron 4           | Splice-site variant | NM_006359: c.584+1G>T | - | Severe mental retardation, microcephaly, seizure, strabismus | Riess et al., 2012 |
| Intron 4           | Splice-site variant | NM_006359: c.584+5G>A | - | Global developmental delay, febrile seizure, delayed myelination, ataxia | Mercimek-Mahmutoglu S, et al. 2015 |
| Intron 5           | Splice-site variant | c.794-2A>G | - | Severe mental retardation, microcephaly, focal seizure, autism | Fung et al. 2017 |
| Intron 6           | Splice-site variant | NM_006359: c.899+3_899+6del | Skipping of exon 6 (mRNA validation) | Mental retardation, microcephaly, hearing impairment, MRI was normal | Zhang et al. 2020 |
| Intron 9           | Splice-site variant | NM_006359: c.1141-8C>A | multiple aberrant transcripts (mRNA validation) | Severe developmental delay, microcephaly, epilepsy, ataxia, verbal language absent, MRI was normal | Ieda et al. 2019 |
| Intron 10          | Splice-site variant | c.1151-1G>A | - | Mental retardation, microcephaly, electrical status epilepticus in sleep, ataxia, cerebellar atrophy, | Zanni et al., 2014 |
| Intron 11          | Splice-site variant | c.1463-1G>A | Skipping of exon 12 (mRNA validation) | Mental retardation, epilepsy, microcephaly, verbal language absent, ataxia, cerebellar atrophy | This report |
According to the previous reports, CS specific brain MRI presented as CA. CA occurred mainly in the inferior parts of vermis and cerebellar hemisphere, especially in the cerebellar vermis. This patient has received two brain MRI scans at the age of 9 month and 3 year, respectively. The first brain MRI showed normal intracranial structures, but the later brain MRI showed the inferior parts of cerebellar vermis dysplasia and the inferior orifice of the fourth ventricle enlargement (Figure 1), which indicated that the cerebellum was slowly atrophy and its further development should be monitored during follow-up. It was reported that the occurrence of CA was mainly related to the extensive progressive loss and degeneration of cerebellar Purkinje cells caused by SLC9A6 gene variant in the CS. The above results indicated that the occurrence of CA was progressive in the CS. According to the neuroimaging data of 17 CS patients, the average age of onset of CA was about 11 years old. Therefore, although the brain MRI scans indicate no abnormality, the possibility of CS should also be considered in the patients with mental retardation less than 1 year of age. Their brain MRI should be monitored constantly and regularly after onset.

There are still some limitation in this study. On the one hand, transcription analysis in the patient was not investigated. It was reported that peripheral blood mRNA analysis is helpful for ascertainment of alternative splicing of SLC9A6 induced CS. On the other side, the correlation between genotype and phenotype is still need further investigation.

In conclusion, few Chinese CS patients have been reported based on the current literature. This case described the clinical manifestations of a Chinese CS patient, including epilepsy, microcephaly, ataxia, and progressive cerebellar atrophy. SLC9A6 NM_001042537.1: c. 1463-1G> A was found to be a pathogenic variant. The novel finding broadens the spectrum of SLC9A6 gene variants in CS and provides a basis for the genetic diagnosis of CS.

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CONFLICT OF INTEREST
The authors declare that they have no competing interests.

DATA AVAILABILITY STATEMENT
The data used to support the findings of this study are included within the article. And the raw data used to support the findings of this study are available from the corresponding author upon request.

ORCID
Yun Xie https://orcid.org/0000-0003-4734-4428

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