Novel ARID1B variant inherited from somatogonadal mosaic mother in siblings with Coffin-Siris syndrome 1

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Abstract. Coffin-Siris syndrome1 (CSS1; Online Mendelian Inheritance in Man no. 135900) is a multiple malformation syndrome characterized by intellectual and/or developmental delay, and hypoplastic or absent fifth fingernails and/or toenails. AT-rich interaction domain-containing protein 1B (ARID1B) is the most frequently mutated gene in CSS1 and the majority of reported cases are sporadic. Using whole-exome sequencing, the present study identified two siblings with CSS1 with a novel heterozygous co-segregating pathogenic variant in the ARID1B gene (c.3468_3471del). Additionally, the current study confirmed a 4% somatic ARID1B mosaicism in the patient's mother. The results expanded the spectrum of known ARID1B pathogenic variants. To the best of our knowledge, the present study is the first to provide experimental evidence that an ARID1B pathogenic variant can be inherited from a clinically healthy somatogonadal mosaic mother.

Introduction

Coffin-Siris syndrome1 [CSS1; Online Mendelian Inheritance in Man (OMIM) no. 135900] was first described as a multiple malformation syndrome in 1970 and is characterized by intellectual and/or developmental delay and hypoplastic or absent fifth fingernails and/or toenails (1-3). Haploinsufficiency of the AT-rich interaction domain-containing protein 1B (ARID1B) is the most frequently mutated gene in CSS1 and the majority of reported cases are sporadic. Using whole-exome sequencing, the present study identified two siblings with CSS1 with a novel heterozygous co-segregating pathogenic variant in the ARID1B gene (c.3468_3471del). Additionally, the current study confirmed a 4% somatic ARID1B mosaicism in the patient's mother. The results expanded the spectrum of known ARID1B pathogenic variants. To the best of our knowledge, the present study is the first to provide experimental evidence that an ARID1B pathogenic variant can be inherited from a clinically healthy somatogonadal mosaic mother.

Mosaicism is a biological phenomenon that describes an individual who has developed from a single fertilized egg and has two or more populations of cells with distinct genotypes (12). Specific types of mosaicism describe the parts of the body that harbor the variant cells and the potential for transmission to offspring, including germline mosaicism (also known as gonadal mosaicism), somatic mosaicism and somatogonadal mosaicism (a combination of germline and somatic mosaicism) (12-16). Postzygotic mosaicism refers to mutations that result in distinct cell populations within the same individual when somatogonadal mosaicism cannot be fully excluded (14-16). The developmental timing and cell lineage are affected and, combined with the phenotypic consequences of the pathogenic variant, ultimately determine
the tissue distribution of mosaicism (somatic, germline or somatogonadal), as well as the patterns of disease reoccurrence within families (12,14). Somatogonadal mosaicism that arises at an early embryonic stage can involve both somatic and germ cells, and individuals with somatogonadal mosaicism are at risk of having affected children (12,14).

The present study reported two siblings with CSS1 and demonstrated rare genetic inheritance from their mother with somatogonadal mosaicism.

Materials and methods

Ethical statement. Ethical approval for the present study was obtained from the Institutional Review Board of the Children’s Hospital of Chongqing Medical University, Chongqing, China (ethical approval no. 2018-64). Written informed consent was obtained from the parents of the siblings.

Clinical characterizations. The patient in the present study was a girl aged 6 years and 3 months, born full-term to non-consanguineous Chinese parents. She was delivered normally (gravida 3; para 3) with a birth weight of 4,150 g. The length of the baby and head circumference at birth were unknown. Feeding problems were observed during the infant period. The first baby of the parents had died shortly following birth.

The patient came to hospital and her mother reported that the patient had intellectual and linguistic developmental delays. The patient spoke her first words ‘mum mum’ at the age of 4 years and had slightly improved; however, she only said ‘mum mum’ during the clinical evaluation. She began to walk by herself at 2 years old. She has a severe intellectual delay and an IQ<32 according to the Wechsler Preschool and Primary Scale of Intelligence test (17). According to the operation manual, the average score is 100 and the 95% confidence interval is 70-130.

Upon examination, the patient weighed 14 kg (z-score, -2.91), her height was 105.2 cm (z-score, -2.23), head circumference was 46.5 cm (~20-month average) and BMI was 12.7 (z-score, -2.05), all of which are abnormal. She displayed dysmorphic features, including brachytelephalangy and hypoplastic nails of the fifth digit phalanges and fingers, coarse facial features, low hairline, short philtrum, thick lips, mild dysphagia, micrognathia, small breasts, and the patient could not walk for moderate scoliosis (Fig. 1A-C) and body hypertrichosis. Brain MRI and metabolic screening were normal.

The patient's older brother exhibited similar disease onset and progression. He was 10 years old at the time of the current study and had learning difficulties. Physical examination demonstrated similar dysmorphic features to the patient, except for moderate scoliosis (Fig. 1B). The parents were clinically healthy and exhibited no dysmorphic features.

Whole-exome sequencing (WES). Peripheral blood samples were collected from the patient and her family in EDTA tubes. The genomic DNA of the patient, her brother and their parents were screened for genetic variations using trio WES. Briefly, the DNA was sheared using an ultrasonic processor (version no. KQ218; Kunshan Ultrasonic Instruments Co., Ltd.) and hybridized with the xGen Exome Research Panel probe sequence capture array (version no. 1.0; Integrated Device Technology, Inc.) to enrich the exonic region, according to the manufacturer's protocol. Quantitative PCR was performed for effective molecular concentration detection of the exome libraries using a KAPA Library Quantification kit (cat. no. KR0405-v8.17; the kit included SYBR-Green; Kapa Biosystems, Roche Diagnostics), according to the manufacturer's protocol. The primers and thermocycling conditions are presented in Table I. The size distribution and concentration of the exome libraries was tested using Bioanalyzer 2100 (Agilent Technologies, Inc.), according to the manufacturer's protocol. A standard curve was generated and was used to convert the average Cq score for each library. The average size-adjusted concentration for each library that was assayed was calculated by multiplying the calculated average concentration using the following formula: Size of DNA standard in bp (452)/average fragment length of library in bp.

Sanger sequencing. In order to validate the pathogenic variants detected by WES, the genomic DNA of the patient, her brother and their parents was evaluated using PCR and the products were analyzed using Sanger sequencing. PCR amplification was performed using a KAPA2G Robust HotStart PCR kit (Kapa Biosystems) on a Hema 9600 PCR Thermo Cycler (Zhuhai Hema Medical Instrument Co., Ltd.), according to the manufacturer's protocol. The primers and thermocycling conditions are presented in Table II.

Sequences were assembled and analyzed using the DNASTAR Laser gene software package (version no. 7.1; DNASTAR, Inc.; www.dnastar.com/documentation).

Amplicon-based deep sequencing. A two-stage PCR was performed. For the first stage of PCR, the PCR reaction system was prepared using sample DNA. Each step of the two-stage PCR was identical to that described previously, except for the specific primers (Chigene; not commercially available). The product directly served as the template for the second stage of PCR with the same procedure, except for the second pair specific primers. The primers used in each stage were specifically designed by Chigene (Chigene Translational Medical Research Center Co. Ltd.; chigene.org) and are not commercially available. The purified DNA products underwent Next Generation Sequencing (NGS)-based ultra-deep sequencing (sequencing depth >500,000X). The sequencing data were then used for bioinformatics analysis.

Bioinformatics analysis. Raw image files were processed using a BCL2FASTQ software package (version no. 1.8.4; Illumina, Inc.; support.illumina.com/content/dam/illumina-support/documents/documentation/software_documentation/bcl2fastq/bcl2fastq_letterbooklet_15038058bpmi.pdf) to generate fastq data. Low-quality variations of the quality score <20 were filtered out. The sequencing reads were aligned to the National Center for Biotechnology Information (www.ncbi.nlm.nih.gov/) human reference genome using Burrows-Wheeler Aligner (version no. 0.7.11-r1034; https://github.com/lh3/bwa). SAMtools (version no. 0.1.19; github.com/samtools/samtools) and Pindel (version no. 0.2.4; github.com/genome/pindel) software were used to call single-nucleotide variants and indels of the reads. The minor allele frequency (MAF) was annotated using the databases dbSNP (version no. 151; http://www.ncbi.nlm.nih.gov/
The present study therefore identified a de novo heterozygous 4-bp deletion (c.3468_3471del) in the 13th exon of the ARID1B gene (NM_020732; Fig. 2). This deletion is predicted to result in a frame shift and premature termination. To the best of our knowledge, this pathogenic variant in the ARID1B gene has not previously been reported. De novo occurrence was confirmed using Sanger sequencing of samples from the patient and her parents (Fig. 2). No other pathogenic variants were identified in screening of all associated CSS genes (ARID1B, SMARCB1, SMARCA4, SMARCE1, ARID2, DPF2, SMARCC2 and SMARCA2).

Amplicon-based deep sequencing. After excluding the possibility of incomplete penetrance of CSS1, one of the parents was suspected to be a gonadal mosaic. In order to evaluate this hypothesis, amplicon-based deep sequencing was performed using DNA from the peripheral blood of the patient’s parents, which confirmed a 4% somatic ARID1B mosaicism in the patient’s mother (Fig. 3). The previous detection of the ARID1B pathogenic variant in the patient’s brother indicated the presence of somatogonadal ARID1B mosaicism in the mother.

Discussion

CSS was first reported by Coffin and Siris (1) and is characterized by notable hypoplasia of the fifth digit phalanges and/or nails, coarse facial features and sparse scalp hair (1). Worldwide, >100 CSS cases with ARID1B pathogenic variants have been reported in the last two years and the clinical features have been summarized (7,8). Previous evidence has indicated that the majority of CSS cases exhibit a certain degree of intellectual and/or developmental delay (4,5,8,19). There are two facial features that have been categorized (4,20). Patients with classical CSS have coarse facial features, bushy eyebrows and thick vermilion of the lips, while patients with variant CSS collectively display a less coarse facial features, thinner eyebrows and thin vermilion border of the lips. However, both CSS and ARID1B-associated disorders have a phenotypic spectrum and cannot always be distinguished as two separate (classical or variant) categories. As a result of this phenotypic heterogeneity, no widely convincing clinical criteria and key diagnostic
Table I. Primers and thermocycling conditions of quantitative-PCR for effective molecular concentration of the exome library.

| Primer    | Sequence                      | Step 1                                      | Step 2                                      | Step 3                                      |
|-----------|-------------------------------|---------------------------------------------|---------------------------------------------|---------------------------------------------|
| Forward   | 5′-AATGATACGGCGACCACCCAGA-3′ | Pre-denaturation at 95°C for 5 min          | Denaturation at 95°C for 30 sec, annealing at 60°C for 45 sec, 35 cycles | Melt curve analysis at 65-95°C              |
| Reverse   | 5′-CAAGCAGAAGACGGCATAAG-3′   |                                             |                                             |                                             |

Table II. Primers and thermocycling conditions used for Sanger sequencing of the AT-rich interaction domain-containing protein 1B gene.

| Primer    | Sequence                      | Step 1                                      | Step 2                                      | Step 3                                      |
|-----------|-------------------------------|---------------------------------------------|---------------------------------------------|---------------------------------------------|
| Forward   | 5′-GCCATCAGCAGGTTCCCTAA-3′   | Predenaturation at 95°C for 5 min           | Denaturation at 95°C for 30 sec, annealing at 60°C for 30 sec, chain extension at 72°C for 30 sec, 30 cycles | Final extension at 72°C for 10 min          |
| Reverse   | 5′-CGCCACTTACAGGAGATGG-3′    |                                             |                                             |                                             |

Figure 2. Pedigree of the family and DNA sequencing showed a heterozygous 4-bp deletion (c.3468_c.3471delCGAA+) in the 13th exon of the ARID1B gene in the patient and her brother; this variant was not present in the parents. ARID1B, AT-rich interaction domain-containing protein 1B.
features have yet been outlined (1,3,5,7,19,20). Thus, molecular
testing currently has an important role in the diagnosis of CSS.

In 2012, CSS was reported to be caused by pathogenic
variants in genes encoding subunits of the SWI/SNF complex,
which function as chromatin remodeling factors (21‑23). These
include SMARCB1, SMARCA2, SMARCA4, SMARCE1,
ARID1A and ARID1B at 22q11.23, 9p24.3, 19p13.2, 17q21.2,
1p36.11 and 6q25.3 (21‑23). Additional genes (including SOX11,
ADNP, PHF6 and TBC1D24 at 2p25.2, 20q13.13, Xq26.2 and
16p13.3,) have also been identified (24). Evidence has indicated
that pathogenic variants in ARID1B are the primary genetic
cause of CSS, accounting for 51‑75% of cases (3,5‑7) in
European, Japan, Canada, USA, between 2013‑2019. However,
ARID1B is the most frequently mutated gene in unspeci
fied intellectual disability cohorts (~1%) from the United
Kingdom and Germany in the past 8 years (7,25,26). ARID1B
pathogenic variants are associated with phenotypes that vary
from non‑syndromic intellectual disability to CSS (7).

The patient and her brother presented with intellectual
disabilities, speech problems, developmental delay, hypo
plastic fifth fingernails and/or toenails and brachytelephalangy.
Additionally, the patient and her brother exhibited dysmorphic
features, including coarse facial features, low hairline, bushy
eyebrows, thick lips, scoliosis, body hypertrichosis and dental
anomalies. The two siblings were clinically diagnosed with
CSS1; this was confirmed using WES, which demonstrated
heterozygous ARID1B variants [c.3468 (exon13)_c.3471 (exon13) delCGAA] present in both siblings.

The majority of reported CSS1 cases are caused by
heterozygous pathogenic variants in the ARID1B gene are
sporadic (5,8). However, exceptions have been noted, including
a pathogenic variant of ARID1B that was passed down from an
affected mother to her son (19). In a consanguineous Emirati
family, a heterozygous variant of ARID1B was present in three
affected siblings; however, this was absent in their parents and
unaffected siblings (8). This indicated that one of the parents
was a gonadal mosaic.

In the present study, the heterozygous pathogenic variant
[c.3468 (exon13)_c.3471 (exon13) delCGAA] was identified
using WES in two siblings; however, this was not initially
observed in the parents. Therefore, it was hypothesized that this
was a de novo pathogenic variant. Additionally, we hypothesized
that one of the patient's parents exhibited gonadal mosaicism for
the pathogenic variant, which has previously been reported (8);
however, this had not been verified by laboratory tests. In
order to test this hypothesis, amplicon‑based deep sequencing
was performed using samples from the patient's parents. An
ARID1B pathogenic variant (4%; chr:157505487‑157505490) was
observed in the mother; however, this was absent in the
father. This indicated that the mother exhibited the somatogonadal mosaic ARID1B variant. To the best of our knowledge,
this is the first report based on experimental evidence of an
ARID1B pathogenic variant in CSS1 that was inherited from
the clinically healthy somatogonadal mosaic mother.

In summary, a heterozygous ARID1B pathogenic variant
[c.3468 (exon13)_c.3471 (exon13) delCGAA] was identified in the patient and her brother, both of whom exhibited the
classical phenotype of CSS1. The results of the present study
expand the spectrum of known ARID1B pathogenic variants,
as well as the ethnic backgrounds of reported cases. To the
best of our knowledge, the present study is the first to report
two siblings with a pathogenic ARID1B variant inherited from
their mother who exhibited somatogonadal mosaicism.

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Availability of data and materials

The datasets used and/or analyzed during the present study are
available from the corresponding author on reasonable request.

Authors' contributions

ZM, CQ and DY conceived and designed the current study
and analyzed data. ZM and DY prepared the manuscript. All
authors read and approved the final manuscript.

Ethics approval and consent to participate

Ethical approval for the present study was obtained from the
Institutional Review Board, Children's Hospital of Chongqing
Medical University, Chongqing, China (approval no. 2018‑64).

Informed consent for publication

Consent for publication of the patient's data/images in this
paper was obtained from the parents.

Competing interests

The authors declare that they have no competing interests.
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