INTRODUCTION

Angelman syndrome (AS) is a neurodevelopmental disorder characterized by severe intellectual disability, lack of speech, ataxia of gait, seizures, a characteristic electroencephalography (EEG), and a unique behavior that includes any combination of happy demeanor, easily excitable personality, frequent laughter, and stereotypes (Bird, 2014; Buiting, 2010). AS is caused by the lack of expression of the maternally inherited UBE3A gene (OMIM 601623) in neurons. Around 10% of patients with a clinical diagnosis of AS (AS-like) do not have an identifiable molecular defect. Some of these patients harbor alternative genetic defects that present overlapping features with AS.
Some of these Angelman-like syndrome patients harbor alternative genetic defects that present overlapping clinical features with AS (Tan, Bird, Thibert, & Williams, 2014).

Genomic approaches such as array comparative genomic hybridization and whole-exome sequencing have already been useful to identify alternative genes responsible for other heterogeneous genetic diseases such as Rett, Kleefstra, and Smith–Magenis syndromes (Berger et al., 2017; Kleefstra et al., 2012; Sajan et al., 2016). Here, we identified with exome sequencing a novel heterozygous SMARCE1 (OMIM 603111) splicing variant in a patient with an Angelman-like phenotype.

2 | MATERIALS AND METHODS

2.1 | Ethical compliance

The protocol for the study has been approved by the institutional Ethics Committee of Institut d’Investigació i Innovació Parc Taulí I3PT and the corresponding informed consent has been obtained from the parents.

2.2 | Patient

The proband is a 14-year-old boy who was born at term to non-consanguineous parents following a normal pregnancy. The

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**FIGURE 1** Molecular characterization of the SMARCE1 (NCBI RefSeq NM_003079.4) c.237+1G>T splicing variant. (a) Sanger sequencing chromatographs showing the SMARCE1 c.237+1G>T variant in the patient but not in his unaffected parents. The variant is indicated by black asterisks. (b) cDNA analysis of the SMARCE1 c.237+1G>T variant in peripheral blood. PCR amplification products of exons 3 to 7 were run in a gel electrophoresis. The patient showed two bands compared to the negative control. Sanger sequencing chromatographs indicate skipping of exon 5. The start of the exon is indicated by a dashed line. (c) Schematic structure of the human SMARCE1 gene and protein illustrating the predicted variant effect on splicing and protein. Exons are shown as boxes and introns as lines. Protein functional domains are shown as boxes. The amino acid deletion is delimited by black lines. Proline-rich domain (Pro-rich) 5–65 aa, High Mobility Group (HMG) 66–134 aa, Coiled-Coil domain 220–319 aa, glutamic acid-rich domain (Glu-rich) 320–411 aa
patient had sucking difficulties during the neonatal period. At the age of 3 months, he suffered from seizures and at 6 months, he was found to be hypotonic. He presented global developmental delay: He sat unsupported at 12 months, walked independently at 27 months, and speaks only three words. Dentition was delayed. At 2 years old, neurological examination detected severe intellectual disability, ataxia of gait, receptive and non-verbal communication skills higher than verbal ones and frequent drooling. He presented an abnormal electroencephalogram (EEG), although it was not the characteristic found in AS patients. The behavioral phenotype included happy demeanor, easily excitable personality, hyperactivity, attention deficit, stereotypies, attraction to water, aggressiveness, and autistic features. A clinical suspicion of AS was raised which was not confirmed molecularly. AS testing included methylation PCR of the 15q11.2-q13 region, \textit{UBE3A} sequencing, and \textit{UBE3A} MLPA analysis (SALSA MLPA P336-A2, MRC Holland, Amsterdam, The Netherlands). In addition, subtelomeric MLPA (SALSA MLPA P070), Autism MLPA (SALSA MLPA P343-C1) and 60K array-based comparative genomic hybridization (aCGH) were performed with normal results.

2.3 | Whole-exome sequencing

Trio whole-exome sequencing of the patient and his parents was performed using the SureSelect Human All Exon V5+UTR kit (Agilent Technologies, Santa Clara, CA, USA). Sequencing was performed on an Illumina Hiseq2000 platform (Illumina, San Diego, CA, USA) producing 2x100nt paired-end reads at the National Centre of Genomic Analysis (CNAG-CRG, Barcelona, Spain). High-quality reads were aligned to the GRCh37 decoy reference genome used by the 1000 genomes project (hs37d5) using the GEM3 aligner and variants identified following GATK Best Practices (DePristo et al., 2011). Coverage was assessed using GATK Depth of Coverage while ignoring reads with mapping quality <20 and bases with base quality <30.

All exome variants were filtered for allele frequencies <0.001 in the ExAC database (Lek et al., 2016), and their predicted impact on the protein (nonsense, frameshift, splice site, and missense variants were prioritized). The final candidate variant was confirmed by direct Sanger sequencing in the patient and excluded in his parents.

2.4 | RNA analysis

RNA was extracted from isolated peripheral blood buffy coat stored at −196°C in liquid nitrogen using the Biostic Blood Total RNA Isolation Kit Sample (MO BIO Laboratories, Inc.), and cDNA synthesis was carried out using the PrimeScript™ RT reagent Kit (Takara Bio Inc.). Primers amplifying the region, including exons 3, 4, 5, 6, and 7 of \textit{SMARCE1} gene (NCBI RefSeq NM_003079.4), were designed in order to analyze mRNA splicing.

3 | RESULTS

Trio whole-exome analysis identified a splice site variant c.237+1G>T in the \textit{SMARCE1} gene (NCBI RefSeq NM_003079.4) after filtering the data according to a
dominant de novo model of inheritance, a population allele frequency of <1/1,000 and a predicted impact on the protein. Variants in the SMARCE1 gene are known to cause Coffin–Siris syndrome (CSS5; Coffin–Siris syndrome 5, OMIM 616938), which is a rare congenital syndrome affecting many organs, characterized by moderate to severe intellectual disability (Kosho & Okamoto, 2014; Santen, Emmelien, Vulto-van Silfhout, Pottinger, & Van Bon, 2013).

The presence of the variant was confirmed by Sanger sequencing in the index patient, whereas the variant was not detected in his unaffected parents (Figure 1a). In order to analyze the effect of the splice site variant on mRNA processing, cDNA analysis was performed on the patient and a control sample. Amplification of exons 3 to 7 resulted in an additional smaller fragment in the patient suggesting exon skipping. Sanger sequencing of the cDNA confirmed the skipping of exon 5 in the patient sample (Figure 1b). Deletion of exon 5 results in an in-frame deletion of 27 amino acids, removing the last part of the Pro-rich (Proline-rich) domain and the start of the HMG (High Mobility Group) domain (Figure 1c), which is essential for the proper functioning of the protein (Lomelí & Castillo-Robles, 2016).

DISCUSSION

Using trio whole-exome sequencing, we identified a novel splice site variant resulting in an in-frame deletion in the SMARCE1 gene in a patient with an AS-like phenotype. Pathogenic variants in SMARCE1 and another five genes (SMARCB1, SMARCA4, SMARCA2, ARID1A, and ARID1B) encoding subunits of the switch/sucrose non-fermenting (SWI/SNF) ATP remodeling complex cause CSS (Miyake, Tsurusaki, & Matsumoto, 2014) which is a rare congenital syndrome characterized by developmental delay, moderate to severe intellectual disability, hypoplastic or absent fifth fingernails or toenails, distinctive facial features, hypertrichosis, sparse scalp hair, and hypotonia (Kosho & Okamoto, 2014).
Germline \textit{SMARCE1} heterozygous loss-of-function variants have been found in young patients with cranial and spinal meningiomas, consistent with a tumor suppressor mechanism (Lomell & Castillo-Robles, 2016; Smith et al., 2014) while missense variants cause Coffin–Siris syndrome (Kosho & Okamoto, 2014).

Here, we describe a splicing variant (c.237+1G>T) in the \textit{SMARCE1} gene leading to an in-frame deletion of 27 amino acids, removing part of the HMG domain (Figure 1c). The patient presents CSS clinical features. No spinal or intracranial meningiomas, which are characteristic of pathogenic loss-of-function variants in \textit{SMARCE1}, were detected in a recent MRI, which showed a dysgenesis and hypoplasia of the corpus callosum and a global dilatation of the ventricular system, characteristic of Coffin–Siris patients. Interestingly, Smith et al., (2013) reported a splicing variant similar to the variant found in our patient, c.237+2T>C. It was identified in two members of a family with multiple spinal meningiomas with no clinical symptoms of CSS. RNA analysis of the affected individuals presented two alternatively spliced \textit{SMARCE1} transcripts, one leading to a premature STOP codon and a less abundant second transcript leading to the same in-frame deletion as that in our patient: In our patient, the c.237+1G>T variant leads only to one alternative splice variant, the 27 amino acid in-frame deletion, suggesting that the abundance of the in-frame deletion transcript leads to a gain of function or dominant negative effect (Tsurusaki et al., 2012) and the appearance of a CSS phenotype.

The patient described here shows almost all the consistent and frequent clinical features associated to AS (Williams, Driscoll, & Dagli, 2010) except for the microcephaly and the frequent laughter/smiling (Table 1). Also, other associated AS features such as attraction to water, a happy demeanor, or frequent drooling are present. This led to the clinical diagnosis of AS which was not confirmed molecularly.

Clinical reevaluation of the patient after the identification of the \textit{SMARCE1} pathogenic variant showed the presence of clinical features associated to CSS but not to AS (Table 1). Among them, the characteristic hypoplastic nail on the 5th finger of the left hand, a coarse facies, sparse scalp hair, hypertrichosis in the back and dysgenesis and hypoplasia of the corpus callosum (Figure 2). CSS patients, like AS patients, present severe developmental delay, speech impairment with expressive language more severely affected than receptive language, moderate to severe intellectual disability and behavioral abnormalities such as hyperactivity and autistic features (Table 1) reflecting the clinical overlap between the two syndromes. However, the patient described here presents some other features characteristic of AS which have not been described in CSS before such as the ataxia of gait and the stereotypes.

To date, only six individuals with \textit{SMARCE1} missense pathogenic variants have been reported (Zarate et al., 2016). Here, we describe the first patient with a pathogenic splicing variant in the CSS gene, \textit{SMARCE1}, who had a diagnosis of AS-like and who presents some clinical features characteristic of AS, which have not been previously associated to CSS. Taking into account these results, we believe that CSS should be added to the expanding list of differential diagnoses for AS, probably accounting for some of the molecularly undiagnosed AS-like patients.

The increasing use of exome sequencing in diagnostic laboratories will allow the analysis of all those genes that are involved in severe neurodevelopment disorders in patients who present AS clinical features, improving the diagnostic rate, and providing knowledge of the phenotypic spectrum of AS-like causative genes, among them those responsible for Coffin–Siris syndrome.

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**CONFLICT OF INTEREST**

The authors declare no conflict of interest.

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