Retrospective Diagnosis of a Novel ACAN Pathogenic Variant in a Family With Short Stature: A Case Report and Review of the Literature

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Short stature is a frequent disorder in the pediatric population and can be caused by multiple factors. In the last few years, the introduction of Next Generation Sequencing (NGS) in the molecular diagnostic workflow led to the discovery of mutations in novel genes causing short stature including heterozygous mutations in ACAN gene. It encodes for aggrecan, a primary proteoglycan component specific for the structure of the cartilage growth plate, articular and intervertebral disc. We report a novel ACAN heterozygous pathogenic variant in a family with idiopathic short stature, early-onset osteoarthritis and osteoarthritis dissecans (SSOAOD). We also performed a literature review summarizing the clinical characteristic of ACAN’s patients. The probands are two Caucasian sisters with a family history of short stature and osteoarthritis dissecans. They showed dysmorphic features such as mild midface hypoplasia, brachydactyly and broad thumbs, especially the great toes. The same phenotype was presented in the mother who had had short stature and suffered from intervertebral disc disease. DNA sequencing identified a heterozygous pathogenic variation (c.4390delG p.Val1464Ter) in the sisters, with a maternal inheritance. The nonsense mutation, located on exon 12, results in premature truncation and presumed loss of protein function. In terms of treatment, our patients underwent recombinant human growth hormone replacement therapy, associated with gonadotropin releasing hormone therapy, in order to block early growth cessation and therefore reach a better final height. Our case suggests that SSOAOD ACAN related should be considered in the differential diagnosis of children with autosomal dominant short stature and family history of joints disease.

Keywords: short stature, ACAN, mutation, aggrecans, osteoarthritis dissecans
INTRODUCTION

Short stature is defined as a height of at least two standard deviations below the average observed in age and sex control population (Hauer et al., 2017). It’s a frequent disorder affecting 3% of the pediatric population and one of the most common causes of pediatric endocrinologist evaluation (Stavber et al., 2020). According to its etiology, short stature can be classified into primary growth disorder, secondary growth disorder and idiopathic short stature (ISS) (Xu et al., 2018). In addition to forms determined by common variants with polygenic inheritance, recent studies have highlighted that monogenic defects could be the cause of the growth disorder observed in non-syndromic children (Grüner and Jorge, 2018; Vasques et al., 2019). More than 700 genes are associated with growth failure, with SHOX haploinsufficiency as the most frequent form, estimated to cause 2–3% of idiopathic short stature (ISS) cases (Fukami et al., 2016).

The introduction of NGS in the molecular diagnostic workflow led to the discovery of mutations in novel genes causing short stature including heterozygous mutations in ACAN gene (ACAN, OMIM:*155760) (Sentchordi-Montané et al., 2018). ACAN is located on chromosome 15q26 and consists of 19 exons, ranging in size from 77 to 4224 bp. It encodes for aggrecan, a primary proteoglycan component specific for the structure of the cartilage growth plate, articular and intervertebral disc (Uchida et al., 2020). Aggrecan is fundamental in articular cartilage, providing the hydrated gel structure necessary for the load-bearing properties of joints (Dateki, 2017) as well as chondrocyte and bone morphogenesis (Quintos et al., 2015). The core protein consists of three globular domains (G1, G2, and G3), interglobular domain (IGD), and centrally located glycosaminoglycan attachment region (GAG), consisting of chondroitin sulfate attachment domains (CS1 and CS2) and keratan sulfate attachment domain (KS) (Figure 1). G1 domain forms interactions with hyaluronan, G2 biological function is still not known whereas the G3 domain binds to different extracellular proteoglycans (i.e., tenascin and fibulin) via its C-type lectin repeat domain (CLD) (Gkourogianni et al., 2017). ACAN biallelic mutations are related to spondyloepimetaaphysseal dysplasia, aggrecan type (SEMD, OMIM#612813) (Tompson et al., 2009), while mutations at the heterozygous state can cause spondyloepiphyseal dysplasia, Kimberley type (SEDK, OMIM#608361) (Gleghorn et al., 2005), short stature associated with or without accelerated bone age (BA), early-onset osteoarthritis (OA) and/or osteoarthritis dissecans (SSOAOD, OMIM#165800) and various idiopathic short stature phenotypes (Stattin et al., 2010; Quintos et al., 2015; Tatsi et al., 2017; Van der Steen et al., 2017).

Here we report two sisters from a family with idiopathic short stature, early-onset osteoarthritis and osteoarthritis dissecans caused by novel heterozygous ACAN variant.

CASE PRESENTATION

The probands are two caucasian sisters (III-1 and III-2, Figures 2A,B), previously referred to another hospital for persistent short stature.

III-1 was born full-term after an uncomplicated pregnancy and delivery. At birth, her weight was 3.3 kg (−0.05 SD), length of 46 cm (−2.12 SD), head circumference of 36 cm (+1.68 SD) and she presented normal psychomotor development (Table 1). Growth rate has been below normal during most of her childhood. At our first evaluation, she was 10.8 years old, her height was 125.8 cm (−2.5 SD) with a genetic target of 150.4 cm (−2 SD). The previous year her physical examination revealed bilateral breast development (Tanner stage B2) and gonadotropin releasing hormone stimulation test was performed (peak FSH value of 18.6 mU/ml, peak LH 7.3 mU/ml). She started Gonadotropin-releasing hormone (GnRH) treatment for precocious puberty with Triptorelin 11.25 mg i.m. every 4 weeks. The father’s height (II-2) was 165.4 cm (−1.84 SD) while maternal height (II-1) was 147.3 cm (−2.6 SD). Maternal family history of short stature was described:
the grandfather’s height (I-2) was 152 cm (−3.9 SD), the grandmother’s height (I-1) was 156 cm (−1.12 SD) and the height of the grandfather’s sister (I-3) was 150 cm (−2 SD) (Figure 2C). They had all the same phenotype characterized by short stature, macrocephaly, short neck, barrel chest, muscle hypertrophy and severe osteoarthritis. III-1 showed dysmorphic features such as macrocephaly, frontal bossing, low-set rotated ears, flat nasal filter, brachydactyly and broad thumbs, especially the great toes. Growth hormone (GH) stimulation test (arginine) produced a peak GH value of 25.6 ng/ml. Additional laboratory and endocrine assessment revealed normal complete blood count, chemistry panel, inflammatory markers, celiac screening, hepatic-renal-thyroid function test, insulin-like growth factor-1 (IGF-1) (Table 1). The hand and wrist radiography showed no
characteristic signs of skeletal dysplasia, other than shortening of the 1st distal phalanx. According to the Greulich and Pyle method, her bone age (BA) was 0.6 years advanced relative to the calendar age (CA). She started recombinant human GH (rhGH) replacement therapy at a dose of 0.25 mg/kg/week when she was 11 years old. At 12.6 years of age, the treatment with GnRH was discontinued and menarche occurred at 14.5 years of age. Moreover, in the last two years, she had persistent pain in both knees: x-ray and magnetic resonance imaging detected multiple osteochondral lesions consistent with osteochondritis dissecans and she underwent a surgical operation. At her latest visit (age 14.7 years) her height was 143 cm (−2.3 SD) with a growth rate of 5.3 cm/year (+1.4 SD).

After starting GH’s therapy, both sisters were seen every 6 months for monitoring their growth and pubertal development but also for undergoing laboratory and endocrine assessment for glucose and lipid metabolism, epatic-renal-thyroid and corticotrope function test. All the evaluations were always normal; the dose adjustment for GH therapy was based on IGF-I levels. No side effects were registered for both GH and GnRH therapy. Clinical, anthropometric, and metabolic features during GH treatment are summarized in Table 2.

The mother had the same phenotype, characterized by short stature, macrocephaly, short neck, frontal bossing, flat nasal filter, brachydactyly and broad thumbs, especially the great toes. Despite having a stature < 2.6 SD, she was never referred to an endocrinologist or didn’t undergo any endocrine assessment. From 35–40 years, she started suffering from severe intervertebral disc disease but didn’t need to perform any surgical procedure.

### METHODS

#### Genetic Analysis

Following written informed consent, the genomic DNA of the patients was extracted from peripheral blood through the ReliaPrep Blood gDNA Miniprep System (Promega), according to the manufacturer's recommendation. The GenetiSure Dx Postnatal Array 4x180K + SNP (Agilent Technologies, USA) was used following standard protocols.

The entire coding region of the SHOX gene (exon 1–exon 6a/6b) and intron–exon boundaries were amplified by PCR and sequenced by Big Dye Terminator Kit (Applied Biosystems, Foster City, CA) through the automatic sequencer ABI PRISM 3100 Genetic Analyzer (Applied Biosystems). Search for deletions/duplications was performed by an MLPA assay using an MLPA Commercial Kit (SALSA MLPA Kit P018-G1 SHOX; MRC-Holland, Amsterdam, Netherlands) following the manufacturer's instructions.

A custom-designed NGS panel including 67 genes [A2ML1, ACAN, ANKRD11, BRAF, CBL, CHD7, COL2A1, COMP, CUL7, FGD1, FGF8, FGFR1, FGFR3, FLNB, GHI, GHR, GHRH, GHRHR, GHSR, GLI2, GLI3, GNAS, HDAC6, HESX1, HMGA2, HRAS, IGFI, IGFIIR, IGFLS, IHH, KAT6B, KDM6A, KRAS, LHX3, LHX4, LIG, LZTR1, MATN3, MEK1, MEK2, NF1, NPPC, NPR2, NRS, OXT2, PCNT, PDE3A, PDE4D, POCI1, POU1F1, PROK2, PROK2, PROP1, PTPN12, RAF1, RIT1, SHH, SHOC2, SLC26A2, SOS1, SOS2, SOX2, SOX3, SOX9, SPRED1, STAT5B, TRIM37] involved in bone growth plate development, hypohalamic-pituitary IGF-1 axis and reported mutated in syndromic short stature was used to investigate the presence of pathogenic variants in the two sibs. An

### Table 1

**Phenotype of our two patients with ACAN mutations.**

| Patient | III-1 | III-2 |
|---------|-------|-------|
| **Birth characteristics** | | |
| Weight (kg) | 3.3 | 3.1 |
| Length (cm) | 46 | 46 |
| HC (cm) | 36 | 35.5 |
| **First visit** | | |
| Age (years) | 10.8 | 7.6 |
| Weight (kg) | 28.9 | 18.6 |
| Height (cm) | 125.8 | 109.6 |
| Target Height | 150.4 | 150.4 |
| Arm Span (cm) | 131.5 | 108 |
| Sit Height (cm) | 131.5 | 108 |
| Arm Span/height (cm) | 0.97 | 0.9 |
| Sit Height/height (cm) | 0.85 | 0.99 |
| **Endocrine evaluation** | | |
| GH Peak at GH stimulation test (ng/ml) | 25.6 | 27.9 |
| IGF-1 (ng/ml) | 275 | 132 |
| **Skeletal system** | | |
| Chronologic age (y) | 10.6 | 7.6 |
| Bone age (Greulich/Pyle) | 11 | 7.6 |

HD, head circumference; GH, growth hormone.
Agilent SureSelectQXT target capture method was utilized with a panel size corresponding to 1.3 Mb. The sequencing probes were designed to cover all coding exons of the 67 genes and the 20 bp flanking sequence from the intron-exon boundary. Sequencing reaction was performed using an Illumina MiSeq and MiSeq v2 300 Cycle Reagent Kits. Variant calling was performed with the SureCall v3.5 software (Agilent Technologies) and VCF files were annotated with the wANNOVAR tool. Finally post aligned, annotated, and categorized sequence data was analyzed using a personalized bioinformatics pipeline: variants were filtered by frequency, excluding those with a MAF ≥0.01 in the public databases: 1000 Genome project (https://www.internationalgenome.org/) and gnomAD (https://gnomad.broadinstitute.org/). We prioritized variants causing frameshift, stop codon, splicing or aminoacidic changes predicted as pathogenic by at least four in silico prediction software (SIFT, Polyphen-2, Mutation Taster, CADD). Classification of variants was evaluated using the consensus guidelines as set out by the American College of Medical Genetics and Genomics (ACMG) guidelines (https://www.acmg.net/ACMG). Sanger sequencing was performed to confirm the presence or absence of these mutations in all the family members.

RESULTS

In the two sisters (III-1 and III-2) the aCGH revealed a normal female Karyotype and SHOX molecular analysis excluded any form of SHOX-haploinsufficiency. A targeted NGS panel containing 67 genes involved in syndromic and non-syndromic short stature detected in both sibs the presence of a heterozygous pathogenic variation in exon 12, namely c.4390delG that causes the changing of the reading frame from GTA encoding Valine at position 464 to stop codon (p.Val1464Ter). The variant, validated by Sanger sequencing, was inherited from the short stature mother (Figure 3).

DISCUSSION AND CONCLUSION

The etiology of short stature is complex and can be related to multiple factors such as GH deficiency, ISS, genetic inheritance, malnutrition, chronic disease (such as hypothyroidism, chronic liver and kidney diseases), constitutional delay of growth and puberty, intrauterine growth retardation, skeletal dysplasia, psychosocial stress and environmental factors (Liang et al., 2020; Wang et al., 2020). Based on new gene sequencing techniques, it is now possible to identify the cause of some short stature cases previously diagnosed as ISS.

We report a novel heterozygous mutation in ACAN (c.4390delG; p.Val1464Ter), located in exon 12, in a family with short stature associated with early-onset osteoarthritis and osteoarthritis dissecans (SOAOD). Nilsson et al. (2014) reported for the first time that heterozygous ACAN mutations could be the cause of short stature and accelerated bone age (Nilsson et al., 2014). Since than an increasing number of ACAN pathogenic variants have been identified. To date, 93 mutations in this gene have been reported in HGMD (http://www.hgmd.cf.ac.uk/ac/index.php), with the majority of the variants (81/93) associated with ISS. Among them, 87.9% were inherited whereas the remaining arose de novo. Up to now 18 different mutations in exon 12 have been described (Figure 1; Table 3) (Gleghorn et al., 2005; Quintos et al., 2015; Gkourogianni et al., 2017; Hattori et al., 2017; Hauer et al., 2017; Tatsi et al., 2017; Van der Steen et al., 2017; Sentchordi-Montané et al., 2018; Zeng et al., 2018; Fukuhara et al., 2019; Uchida et al., 2020; Lin et al., 2021). This represents the exon with the highest number of mutations, accounting for the 19% of ACAN alterations (Liang et al., 2020; Lin et al., 2021). Exon 12 encodes for the chondroitin sulfate attachment domains (CS1 and CS2) and along with exon 11, for the keratan sulfate attachment domain (KS). Thanks to the sulfation of these domains (KS, CS1 and CS2) and the aggregation with hyaluronan, a fixed negative charge is generated (Dateki, 2017). This process is fundamental to cartilage because it attracts ions and water molecules, letting the cartilage bear the high mechanical load found in the skeletal joint (Gibson)
All heterozygous aggrecan mutations known so far cause short stature of similar severity, indicating that they similarly affect growth plate cartilage. Nevertheless, not all of them cause involve the articular cartilage and, therefore, determine early-onset osteoarthritis. Early-frameshift or splice-site mutations are likely to cause early truncation of the protein and may therefore functionally represent haploinsufficiency of ACAN. Patients with these types of mutations have short stature but no evidence of early-onset osteoarthritis, suggesting that ACAN haploinsufficiency primarily affects growth plate cartilage. Previously reported familial osteochondritis dissecans family and our family have missense or nonsense mutations in the C-type lectin domain and articular disease, suggesting that the presence of aggrecan protein with a dysfunctional C-type lectin domain causes a more severe phenotype affecting both growth plate and articular cartilage function (Nilsson et al., 2014). The nonsense mutation found in the present patients results in premature truncation and presumed loss of protein function. Ten frameshift variants, four missense variants and four nonsense variants (Table 3) had been previously identified in exon 12, causing a premature termination codon and the loss of the C-terminal globular domains (G3) (Figure 1; Table 3). G3 is fundamental for the proper protein function, contributing to the interaction between the aggrecan molecule and various components of the extracellular matrix (ECM) (Aspberg, 2012; Quintos et al., 2015). Nevertheless, numerous authors reiterate the importance of performing additional studies to determine whether these variants cause mRNA nonsense-mediated decay (NMD) or allow translation of a truncated aggrecan protein into the cartilage extracellular matrix (Gibson and Briggs, 2016; Stavber et al., 2020). To date, the few functional studies on the consequences of mutations in the G3 domain conducted showed the secretion of a truncated aggrecan, ruining cartilage structure and function with a dominant-negative effect (Tompson et al., 2009; Stattin et al., 2010; Gibson and Briggs, 2016).

The heterozygous variant detected in III-1 and III-2 was maternally inherited and their similar phenotype is consistent with that described in previous reports (Table 3), with short stature below −2.4/−2.5 SD, prognathia, flat nasal bridge, midfacial hypoplasia, posteriorly rotated ears, broad forehead, relative macrocephaly, brachyactyly, broad great toes, short thumbs, lordosis, genu valgum, joint problems (Dateki, 2017; Dateki et al., 2017; Hu et al., 2017; Van der Steen et al., 2017; Kim et al., 2020; Nilsson, 2020). The same characteristics can be detected, also, among patients carrying ACAN mutations located on other domains, underlying the absence of a genotype-phenotype correlation. The three most frequent features described among all ACAN’s patients were short neck, mild midface hypoplasia (frontal bossing, flat nasal bridge, and long philtrum), and brachyactyly, secondly osteoarthritis, spine malformation, short limbs, thoracic deformity (Liang et al., 2020). Both the sibs developed early-onset osteoarthritis (II-1 in the knee, while the III-2 in the ankle) and, subsequently, osteochondritis dissecans. While these clinical presentations are relatively common in patients carrying ACAN mutations, in the other cases with truncating mutations in the G3 domain there

**FIGURE 3** | Genetic analysis. (A) NGS file and (B,C) Sanger sequencing chromatograms reporting the wild-type (B) and the c.4390delG variant identified in the patients (C) in exon 12 (NM_013227) of the ACAN gene.
**TABLE 3 | Clinical findings of patients with ACAN mutation in exon 12 in this study and the literatures.**

| Subject | Mutation | Protein | Type       | Sex/Age | Height (SDS) | Phenotype | Patterns of inheritance/affected family members (n) | References                          |
|---------|----------|---------|------------|----------|--------------|-----------|-------------------------------------------------|-------------------------------------|
| 1       | c.3986dupC | p.Gly1330fs*221 | Frameshift  | NA       | NA           | SEDK      | NA                                              | Gleghorn et al. (2005)              |
| 2       | c.4657G>T  | p.Glu1553* | Nonsense   | F/NA     | −3.20        | ISS, midface hypoplasia, flat nasal bridge, short thumbs, OA, familiar short stature with intervertebral disc disease | Mother/(7)                        |
| 3       | c.4762_4765del | p.Gly1588fs*26 | Frameshift  | M/12.3 yr | −2.70        | ISS, midface hypoplasia, broad great toes, posteriorly rotated ears, familiar short stature with hip joint problems, advanteg BA (13.3 yr) | Father/(2)                        |
| 4a      | c.5391delG | p.Gly1797fs*52 | Frameshift  | M/5.6 yr | −2.80        | ISS, midface hypoplasia, flat nasal bridge, familiar short stature with hip joint problems, advanteg BA (8 yr) | Mother/(3)                        |
| 4b      | c.5391delG | p.Gly1797fs*52 | Frameshift  | M/NA     | −2.60        | ISS, midface hypoplasia, flat nasal bridge, short thumbs, markedly advanteg BA, familiar short stature | Mother/(3)                        |
| 5       | c.5597C>A  | p.Ser1866* | Nonsense   | M/NA     | −2.00        | ISS, prominent forehead, short neck, barrel-shaped chest, brachydactyly, advanteg BA | De novo mutation                   |
| 6       | c.2218A>T  | p.Thr740Ser | Missense   | M/3.0 yr | −3.20        | ISS, frontal bossing, midface hypoplasia, high arched palate, triangular face, brachydactyly, BA equal to CA, familiar short stature | NA/(1)                            |
| 7       | c.2369C>G  | p.Ser790* | Nonsense   | M/14.5 yr | −2.20        | ISS, frontal bossing, broad nose and philtrum, hypertelorism, brachydactyly, hyperlordosis, advanteg BA (16.5 yr), no familiar short stature, | NA/(0)                            |
| 8       | c.6142C>G  | p.Pro2048Ala | Missense   | F/12.5 yr | −2.20        | ISS, brachydactyly, BA equal to CA, familiar short stature | NA/(1)                            |
| 9       | c.463delT  | p.Leu1545Profs*11 | Frameshift  | F/7.8 yr | −3.60        | ISS, midface hypoplasia, short left 4th toe, broad 5th toes, short 1st distal phalanx, advanteg BA (8.10 yr), familiar short stature with early-onset invertebral disc disease | Mother/(3)                        |
| 10      | c.2535_2536insTTCA | p.Pro846Phafs*9 | Frameshift  | M/4.9 yr | −2.47        | ISS, BA equal to CA, no familiar short stature | De novo mutation                   |

(Continued)
| Subject | Mutation | Protein | Type           | Sex/Age | Height (SDS) | Phenotype                                                                 | Patterns of inheritance/affected family members (n) | References |
|---------|----------|---------|----------------|---------|-------------|---------------------------------------------------------------------------|-----------------------------------------------------|------------|
| 11      | c.6404delC | p.Ala2135Aspfs | Frameshift | M/6.9 yr | -3.20       | Disproportionate short stature, midface hypoplasia, advanteg BA (9.9 yr), no familiar short stature | De novo mutation | Tatsi et al. (2017) |
| 12      | c.4852C>T | p.Gln1618* | Nonsense     | M/11.7 yr | -4.00       | ISS, attention deficit hyperactivity disorder, mild dysmorphic facial features as retrognathia and midface hypoplasia, BA equal to CA, familiar short stature with OA and invertebral disc disease | Father/(12) | Tatsi et al. (2017) |
| 13a     | c.6193delC | p.Gln2065Serfs*27 | Frameshift | M/9.10 yr | NA          | ISS, familiar short stature                                               | Father/(3) | Zeng et al. (2018) |
| 13b     | c.6193delC | p.Gln2065Serfs*27 | Frameshift | M/7 yr | NA          | ISS, familiar short stature                                               | Father/(3) | Zeng et al. (2018) |
| 14      | c.5443delC | p.Leu1815fs | Frameshift | M/4 yr | -4.38       | ISS, BA equal to CA                                                       | NA | Lin et al. (2021) |
| 15      | c.5579delC | p.Gly1861fs | Frameshift | F/9.4 yr | -2.91       | ISS, advanteg BA (10.7 yr),                                               | NA | Lin et al. (2021) |
| 16      | c.4138G>T and c.5061T>A | p_Val1380Phe and p.Ser1687Arg | Missense | M/ 45 yr | -9.1       | SEMD, diagnosis of osteogenesis imperfecta at 6 years of age               | NA | Fukuhara et al. (2019) |
| 17      | c.2541del | p.Val848fs | Frameshift | NA | NA          | ISS, OA, OCD                                                              | NA | - |
| 18      | c.5546G>A | p.Gly1849Asp | Missense | NA | NA          | ISS, macrocephaly, frontal bossing, low-set rotated ears, flat nasal filter, brachydactyly and broad thumbs, OA, OCD, familiar short stature with early-onset invertebral disc disease, BA equal to CA | Mother/(3) | Ill-1 (current study) |
| 19a     | c.4390delG | p.Val1464Ter | Nonsense | F/10.8 yr | -2.50       | ISS, macrocephaly, frontal bossing, low-set rotated ears, flat nasal filter, brachydactyly and broad thumbs, OA, OCD, familiar short stature with early-onset invertebral disc disease, BA equal to CA | Mother/(3) | Ill-2 (current study) |
| 19b     | c.4390delG | p.Val1464Ter | Nonsense | F/7.6 yr | -2.40       | ISS, macrocephaly, frontal bossing, low-set rotated ears, flat nasal filter, brachydactyly and broad thumbs, OA, OCD, familiar short stature with early-onset invertebral disc disease, BA equal to CA | Mother/(3) | Ill-2 (current study) |

NA, not available; Yr, year; SEDK, spondyloepiphyseal dysplasia Kimberley type; ISS, idiopathic short stature; OA, osteoarthritis; OCD, osteochondritis dissecans; SEMD, spondyloepimeta physeal dysplasia, aggrecan type.
were only two cases of early-onset osteoarthritis (Gleghorn et al., 2005; Gkourogianni et al., 2017) but no other osteochondritis dissecans. The two sibs presented joint problems at a young age (between 10–12 years), in contrast to previous reports where the common start was late adolescence or even later (Gkourogianni et al., 2017). Their affected mother (II-1) suffered from severe intervertebral disc disease, starting at 35–40 years, indicating wide phenotype variability within this family.

In terms of treatment, although the two sisters didn’t develop precocious puberty, they underwent gonadotropin releasing hormone (GnRH) therapy, to block early growth cessation and therefore reached a better final height. Several studies evaluated the effectiveness of combined GH and GnRH analog treatment for achieving a proper final height in patients with ACAN mutations (Dateki, 2017; Van der Steen et al., 2017; Zeng et al., 2018; Stavber et al., 2020). Gkourogianni et al. (2017) found that the mean height SD for ACAN adult patients previously treated with GH was −2.5, while that of untreated adult individuals was −3.0 (Gkourogianni et al., 2017). Furthermore, Van der Steen et al. (2017) reported that GH-treated adult individuals, who received in addition GnRH analog treatment for two years from puberty (mean age of GnRH treatment’s start between 10–12 years, with normal timing of puberty), gain 5–8 cm at their adult height compared to their family members, with the same sex and ACAN mutation (Van der Steen et al., 2017). Stavber et al. (2020) described the cases of three children who, in addition to GH treatment, received treatment with GnRHa during puberty after 10 years of age. During the combined therapy there was a significant improvement in SD height and growth velocity increased significantly. Nevertheless, the author underlines that it’s not possible to establish the efficacy of the therapy since none of the patients have already achieved the final height at the time of publication (Stavber et al., 2020). Therefore, it is still controversial if ACAN patients need to undergo GH therapy with or without GnRHa to achieve a proper final height and further studies are needed (Dateki, 2017).

In conclusion we identified a novel heterozygous ACAN mutation causing SSOAOD in the family. SSOAOD ACAN related should be considered in the differential diagnosis of children with autosomal dominant short stature and family history of joints disease, even in patients in which BA is equal to CA. Diagnosis of the condition is difficult, and genetic testing may help. As for treatment, growth therapies could be beneficial for improving affected individuals’ adult height.

DATA AVAILABILITY STATEMENT

The datasets for this article are not publicly available due to concerns regarding participant/patient anonymity. Requests to access the datasets should be directed to the corresponding author.

AUTHOR CONTRIBUTIONS

VM collected the anamnestic as well as the biochemical data for the patients. VM performed the literature search, reviewed and extracted data from the papers. MG and SM performed the genetic analysis. VM performed the figures and table designing and the manuscript writing with the assistance of MG, SB, and FP. All authors discussed the results and contributed to the final manuscript.

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