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ARTICLE in EUROPEAN JOURNAL OF MEDICAL GENETICS · JULY 2009
Impact Factor: 1.49 · DOI: 10.1016/j.ejmg.2009.06.001 · Source: PubMed

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Familial Sotos syndrome caused by a novel missense mutation, C2175S, in NSD1 and associated with normal intelligence, insulin dependent diabetes, bronchial asthma, and lipedema

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\textbf{A R T I C L E   I N F O}

Article history:
Received 27 April 2009
Accepted 7 June 2009
Available online 21 June 2009

Keywords:
Sotos syndrome
Vertical transmission
NSD1 mutation
Insulin dependent diabetes mellitus

\textbf{A B S T R A C T}

We report a familial Sotos syndrome in two children, boy and girl, aged 17 and 8 years, and in their 44 year old mother, who displayed normal intelligence at adult age, but suffered from insulin dependent diabetes mellitus, bronchial asthma, and severe lipedema. The underlying missense mutation, C2175S, occurred in a conserved segment of the NSD1 gene. Our findings confirm that familial cases of SS are more likely to carry missense mutations. This case report may prove useful to avoid underestimation of the recurrence rate of SS, and to demonstrate that the developmental delay may normalize, enabling an independent life and having an own family.

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1. Introduction

Sotos syndrome (SS, OMIM 117550) is characterized by pre- and postnatal overgrowth, macrocephaly, typical facial gestalt, large hands and feet, accelerated skeletal age, and developmental delay. Since the first report of this condition in 1964, the diagnosis relied mainly on the clinical phenotype. In 2002, mutations and submicroscopic deletions in the NSD1 gene at chromosome 5q35 were found in 24 of 42 SS patients [1], enabling molecular genetic diagnosis. Some 50–75% of patients demonstrate mutations in the NSD1 gene or a microdeletion at chromosome 5q35 [1–4]. NSD1 encodes the nuclear receptor-binding SET [Su(var)3-9, Enhancer-of-zeste, Trithorax] domain protein 1, which may act as a basic transcriptional factor and bifunctional transcriptional regulator [5]. SET-domain containing proteins (histone lysine methyltransferases) can modify histones and thereby affect gene expression [6].

Today, the Sotos and Weaver syndromes are again thought of as different disorders [4]. For some time it had been proposed that NSD1 mutations may also cause the overgrowth syndrome of Weaver (OMIM 277590), which is less well defined and much rarer than SS, with only some 20 reported cases to date. Sotos and Weaver syndromes show overlapping phenotypes, especially at young ages [4,7]. Phenotypic differences include a slightly different facial appearance and more conspicuous contractures in Weaver syndrome and an advanced dental maturation in SS that is rarely observed in Weaver syndrome [7]. In addition, SS has been associated with a slight increased risk for cancer (in particular Wilms tumor), whereas the Weaver syndrome usually is not, although one case of neuroblastoma was reported in Weaver syndrome [7]. When molecular diagnosis became available, it turned out that classic Weaver syndrome is not caused by NSD1 mutations [2–4].

SS has been associated with a low recurrence risk in sibships and with reduced reproductive fitness. Familial inheritance of SS was first described in 1976 [8]. Since 2003, 16 families have been reported [4,9–11]. In 15 of these families, an NSD1 mutation or 5q35 microdeletion segregating with the disorder was identified, including seven missense [4,10], four frameshift [4,9], two nonsense [4], and one splice donor site mutation [11]. In one family, the causative mutation was not detected [4]. Familial 5q35 microdeletions were observed in monozygotic twin boys only, but not in families with vertical transmission [4]. Missense mutations appear to be more frequent in familial cases than in sporadic cases [4].

Here we report on a family with SS caused by a novel NSD1 missense mutation, C2175S. The affected siblings and their likewise affected mother presented with typical facial dysmorphism, macrocephaly, and tall stature. The siblings had characteristic heart defects. The mother had received estrogen/gestagen treatment of
2. Subjects

The family was of German origin. Patient 3 and her children (patients 1 and 2) had SS, whereas her husband, parents, brother and nephew were unaffected and healthy. Patient 3 grew up in a family providing excellent support in childhood. Her mother was 170 cm in height (+0.5 SD). Her father had been approximately 180 cm tall (+0.4 SD), German shoe size 46 (UK size 11, US men 11.5–12). He was well-educated and a master butcher in his own shop, but had died already in the 1970s of a horseback riding accident. Her brother was a successful trained retail salesman and approximately 198 cm tall (+3 SD), indicating familial tall stature.

2.1. Patient 1

The 10 year old boy (Fig. 1A) was referred to genetic examination by the community pediatrician. He had been delivered by cesarean section at week 36 of gestation after cardiotocogram (CTG) changes had been noticed. Birth weight was 3800 g (+2 SD, 98th percentile) and length 49 cm (+1 SD). His neonatal course was remarkable for prolonged jaundice. An echocardiography revealed an atrioseptal defect (ASD) II. Postnatal growth was accelerated. At age 10 \(\frac{8}{12}\) years, he had proportionate tall stature (169 cm [+3.5 SD]); maternal and paternal height were 180 cm each. He had macrocephaly (occipitofrontal circumference [OFC] 57 cm [+2.4 SD]), high frontal hairline, coarse facial features, prominent mandible, high palate, funnel chest, diastasis of the abdominal wall, and large hands and feet (German shoe size 45 [UK size 10.5, US men 11]). Medical problems included mild myopia and allergic rhinoconjunctivitis (hay fever).

Development had been delayed for the first 9 years of his life, with muscular hypotonia and limited gross and fine motor skills, which his mother described as clumsiness. He received physiotherapy from age 3 months and walked at age 23–24 months. After infancy, he had occupational therapy. Intelligence testing at age 6 years indicated an IQ of 100 in the verbal subtests, and 85 in the performance subtests (Wechsler Intelligence Scale for Children, German Adaptation). He had expressive language delay and received special support and reportedly had been “slow” in elementary school, but then she graduated basic secondary school (Hauptschule) and successfully completed a pastry cook apprenticeship. She had hay fever and bronchial asthma. At the age of 25 years she suddenly felt extremely thirsty and tired, and rapidly lost weight, 10 kg in 4 months. A type 1 diabetes was diagnosed and insulin treatment was initiated. One month later, she became pregnant with her son. In the interval between her two pregnancies, she took anticonvulsives (carbamazepine and valproic acid) for 2 or 3 years because of a suspected epilepsy which was not confirmed later on. When last seen at the age of 44 years, she reported serious medical problems with her diabetes that was difficult to control. She had asthma treated by formoterol inhalation therapy, and lipedema with painful column-like legs and massive accumulations of fat and fluid under the skin of the legs and ankles.

3. Cytogenetic and molecular analyses

Following detailed genetic counselling, both parents of patients 1 and 2 provided informed consent for genetic analyses. Conventional chromosome analysis (Giemsa banding) on cultured blood lymphocytes of patients 1 and 2 revealed normal 46,X and 46,XG karyotypes, respectively. NSD1 microdeletions were ruled out by fluorescent in situ hybridization (FISH) with BAC RP11–118M12, which maps to the critical interval at chromosome 5q35 [1].

Molecular analyses were performed using blood samples from patients 1–3 and from the mother and the brother of patient 3. Genomic DNAs were isolated by a standard salting out procedure. The entire NSD1 protein-coding region (exons 2–23) was amplified using 31 primer pairs, as described previously [3]. PCR fragments were bidirectionally sequenced using the CEQ™ DTCS Quick Start Kit (Beckman Coulter, Krefeld, Germany) and a Beckman CEQ 8000 Genetic Analysis System.

Two heterozygous alterations were found in the family, both in exon 23; a pathologial missense mutation (c.6523T > A, resulting in p.C2175S) (Fig. 2) in all three patients; and a previously reported polymorphism (c.7636G > A, resulting in p.A2546T) [12] in patient 3 and in her mother, but not in patients 1 and 2. The c.6523T > A mutation predicts the exchange of cysteine 2175 to serine in a conserved Cys/His-rich region adjacent to the plant homeo-domain (PHD)-V region which may correspond to another zinc finger-like motif. This cysteine residue is conserved not only in mouse Nsd1 but also in the two known human NSD1 paralogues,
Fig. 1. Facial appearance (A) of patient 1 aged 10 5/12 years, (B, C) of patient 2 aged 8 6/12 years, and (D, E) of patient 3 aged 36 years, respectively. Note the high frontal hairline and mandibular prognathism as typical signs of the Sotos syndrome.
NSD2 and NSD3. The c.6523T > A mutation was not found in the mother and brother of patient 3; a sample of her father was not available. The segregation of polymorphism c.7636G > A, which is present in patient 3 and her unaffected mother, but not in patients 1 and 2 indicates that mutation c.6523T > A occurred de novo on the paternally inherited allele of patient 3.

4. Discussion

Familial cases of SS are rare, the majority of cases are sporadic. The two siblings and their mother described here presented with overgrowth and characteristic dysmorphic signs (including macrocephaly, high frontal hairline, coarse facial features, and prominent pointed chin). Genetic testing identified a pathological NSD1 missense mutation in exon 23 (p.C2175S) in all affected family members. Although the C2175S mutation was not described previously, it is evidently disease-causing because of its position in a conserved domain of the protein. Because a maternally inherited SNP (c.7636G > A) in the same exon (exon 23) did not segregate with the C2175S mutation, the pathogenic mutation must have arisen in the paternally inherited NSD1 allele of patient 3. Our case supports the observation that the diagnosis of familial SS is usually made via an affected child.

Why are some cases of SS heritable, and what are the predisposing factors? Previous studies [2,4] suggested that familial SS is usually transmitted through the mother, accounting for 75% (12/16) of inherited cases. Here, we present another case of maternal transmission. Our report supports the view that missense mutations – in particular, missense mutations outside the SET domain (position 1941–2063) and the zinc fingers – are more likely to be inherited than truncating mutations or microdeletions [4,10]. Furthermore, our data provide evidence that a subset of individuals with missense mutations may show less severe mental impairment than patients with null mutations. The mutation type seems to play a decisive role in reproductive fitness. Patients with NSD1 missense mutations outside the SET domain and the zinc fingers may be milder affected and, therefore, demonstrate higher vertical transmission rates.

Höglund et al. [9] described that mental retardation or physical impairment, if present, may affect reproductive fitness as an independent factor. Patient 3 showed poor coordination and expressive language delay during childhood, but normal development at later ages, enabling a normal professional and family life as an adult. Similarly, the motor and language delays of her son (patient 1) decreased during his first years of life. With speech therapy in the first years, he could attend normal school classes enabling a normal professional development. These observations confirm that especially in familial cases of SS, the initial hypotonia and developmental delay may improve during school age. This increases the likelihood of transmitting the mutation to the next generation. Thus, the risk for affected offspring resembles that of other autosomal dominant disorders [9]. Our report underlines the necessity of molecular studies also in the parents and siblings of patients with SS.

Which type of mutation was identified in this family? The 6523T > A mutation in exon 23 of the NSD1 gene changes the cysteine residue at position 2175 to serine. This finding is in agreement with the fact that all missense mutations that have been described in Sotos syndrome so far are clustered in conserved functional domains in exons 13, 14, 16, 18, 19, 20, 22, and 23, and that missense mutations outside these domains do not cause SS [2–4]. The clinical findings in our family are consistent with the hypothesis that missense mutations may be associated with a milder outcome in a subset of cases. However, the physical features in our patients are relatively prominent, including marked overgrowth, macrocephaly, and cardiac malformation.

Should the diabetes in patient 3 be considered a chance coincidence, or a rare feature of the SS? To our knowledge, there have been no previous reports on SS and diabetes. Overgrowth and an increased tumor risk are well known features of SS, and one might speculate that both problems are caused by malfunction of growth factors. Abnormal expression of growth factors could increase carbohydrate resistance and thus be diabetogenic, and SS has been associated with endocrine and paracrine alterations in the insulin like growth factor (IGF) system [13]. However, patient 3 was diagnosed with insulin dependent (type 1) diabetes mellitus, which is typically caused by an autoimmune disorder, and with bronchial asthma, representing another immunologically mediated disease. She also had lipedema, the causes of which are unknown. Further observations are needed to understand the possible links between SS and diabetes, asthma, and/or lipedema.

Acknowledgements

We thank the participating family members for their help and support.

References


