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Two novel mutations in \textit{RNU4ATAC} in two siblings with an atypical mild phenotype of microcephalic osteodysplastic primordial dwarfism type 1

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List of key features

Dwarfism
Foetal growth retardation
Microcephaly
Mutation
Nucleotides
Osteochondrodysplasias
RNA, small nuclear
Spliceosomes
Syndrome
Microcephalic osteodysplastic primordial dwarfism, type 1

Introduction

Taybi–Linder syndrome or microcephalic osteodysplastic primordial dwarfism type 1 (MOPD1) (MIM # 210710) is a rare autosomal recessive developmental disorder, originally described in 1967 (Taybi and Linder, 1967). The patients present with severe intrauterine and postnatal growth retardation, microcephaly, facial dysmorphism, sparse thin hair and dry skin (Meinecke and Passarge, 1991). Radiological findings include dysplasia of the skeleton with cleft vertebral arches, horizontal acetabula and short and bowed long bones (Sigaudy \textit{et al.}, 1998). Neurological findings typically include dysplasia of the skeleton and cleft vertebral arches, horizontal acetabula and short and bowed long bones (Sigaudy \textit{et al.}, 1998). Neurological findings typically include profound developmental delay, blindness, hearing deficits, central nervous system malformations, early-onset epilepsy and neuroendocrine dysfunction (Pierce and Morse, 2012).

MOPD1 has been shown to result from biallelic mutations in the \textit{RNU4ATAC} gene encoding the small nuclear RNA (snRNA) U4atac, which is a component of the minor spliceosome. Although accounting for splicing of only about 800 introns, the minor spliceosome is involved in the correct splicing of many essential gene products.

Thus, minor intron splicing has a critical role in human development (He \textit{et al.}, 2011).

At present, only around 40 patients with MOPD1 and 10 different \textit{RNU4ATAC} mutations have been reported according to the Human Gene Mutation Database. The condition is usually severe, and the patients do not generally live beyond the age of 3 years (Meinecke and Passarge, 1991). A few cases with a slightly milder phenotype have been reported (Abdel-Salam \textit{et al.}, 2012; Nagy \textit{et al.}, 2012), but no patients have yet been reported to survive into adulthood.

We report on two adult siblings with MOPD1 presenting with an atypical mild phenotypic appearance compared with the previously reported cases.

Clinical reports

The two siblings are the second and third children of healthy nonconsanguineous white parents and have an unremarkable family history. They both presented with prenatal and postnatal growth retardation, microcephaly, developmental delay, cataract, hearing loss and dysmorphic features. Before the establishment of the diagnosis, the cases were reported as unsolved cases (Hansen \textit{et al.}, 2009).

Case 1

A girl, now age 24 years, was born at 38 weeks of gestational age with a birth weight of 1950 g (−3 SD), a length of 43 cm (−4 SD) and a head circumference of 29 cm (−5 SD). She had neonatal hypoglycaemia, which resolved after treatment. In childhood, her skin was affected by severe atopic dermatitis and she had allergies towards egg, milk, nuts and grass. She had several pulmonary infections in early childhood and asthma until 10 years of age. The dysmorphic features included receding forehead, large prominent eyes, arched eyebrows, hypoplasia of the ala nasi, micrognathia, thin hair, small low-set ears and short neck (Fig. 1a and b). Dental examination revealed malocclusion, crowded teeth and...
microdontia with enamel abnormalities. She had tapering fingers and her fingers and toes were broad and short, the skin was dry and the nails were dystrophic (Fig. 1c and d). Radiographs of the long bones, at the age of 1 and 8 years, displayed generalized shortening with metaphyseal broadening and delayed bone age. Ophthalmological examination revealed bilateral cataracts, which were operated at the age of 5 years, and tapetoretinal degeneration. She had menarche at 16 years of age and had normal periods. From the age of 14 years, she had progressive sensorineural hearing loss of 60 dB, partially corrected by hearing aids. At the age of 23 years, she developed severe pneumonia complicated by haemolytic uraemic syndrome and required respiratory support and dialysis for 1 week. Cranial MRI at the age of 24 years demonstrated microcephaly, partial agenesis of corpus callosum and general atrophy. She did not develop epilepsy. At the most recent evaluation, at the age of 24 years, she was severely growth retarded with a height of 142 cm (−5 SD), weight of 35 kg (−6 SD) and a head circumference of 45 cm (−10 SD). Intellectual disability was evident with an IQ of 56, but she was able to live in her own apartment with some support. She had reading skills comparable with a 9-year-old and very limited mathematics skills. She had a slight kyphosis, but no specific orthopaedic problems and was able to walk 5 km.

Case 2
A boy, now aged 17 years, presented like his sister with severe prenatal growth retardation and was delivered at
32 weeks of gestation by Caesarean section with a birth weight of 1079 g (−4 SD) and length of 38 cm (−4 SD). He had atopic dermatitis and allergies in childhood. He was operated on for cryptorchidism. He developed bilateral cataract and received artificial lenses at the age of 4 years, and ophthalmological examination has also revealed tapetoretinal degeneration. He had asthma until 10 years of age. From the age of 10 years, he developed progressive sensorineural hearing loss relieved by hearing aids. The dysmorphic features included receding forehead, bulbous nose, hypoplasia of the ala nasi, full lips, small ears and mild micrognathia (Fig. 2a and b). Dental examination revealed malocclusion and small, crowded teeth with enamel abnormalities. He had tapering fingers, which were broad and short. He had flat feet, syndactyly of second and third toes and dry skin (Fig. 2c and d). Skeletal radiological examination at the age of 12 years (Fig. 3a and b) showed shortening of the long bones, metaphyseal broadening, but otherwise relatively normal configuration. He reached puberty at the age of 15 years. He did not develop epilepsy. At the most recent evaluation, at the age of 17 years, he was severely growth retarded with a height of 139 cm (−7 SD), weight of 30 kg (−6.5 SD) and a head circumference of 46.9 cm (−8 SD). He lived in his parents’ home, attended special school and had reading capability corresponding to...
8 years of age, and no mathematics skills. He had good motor skills, could ride a bicycle, walk 5–10 km and run 3 km without breaks.

**Mutation analysis**

The study was approved by the Scottish Multicentre Research Ethics Committee (04:MRE00/19). Genomic DNA samples from the patients, parents and unaffected sister were analysed at the Institute of Genetics and Molecular Medicine, University of Edinburgh, UK. The RNU4ATAC gene was screened by bidirectional Sanger sequencing, and analyses were performed using Mutation Surveyor (Softgenetics Inc., Pennsylvania, USA). The findings were validated by bidirectional Sanger sequencing at the Department of Clinical Genetics, Odense University Hospital, using SeqMan Pro v.12.0, DNA Star (Wisconsin, USA).

The affected siblings were compound heterozygous for a n.40C>T nucleotide substitution and an 85-base tandem duplication (bp 16–100) in RNU4ATAC, which results in an insertion of an 85-base-pair-long sequence in position n.101.

The parents were both heterozygous for each one of these mutations, and the oldest sister, who was unaffected, was heterozygous for the n.40C>T mutation.

Neither of these mutations have been previously reported in the literature in relation to MOPD1. The n.40C>T was reported in a single individual from south Asia in the Exome Aggregation Consortium corresponding to a population frequency of 0.0093%. No homozygotes for the mutation have been detected.

Figure 4 displays the normal configuration of the U4atac snRNA and localization of the two mutations. The n.40C>T mutation is predicted to disrupt the essential 5’ stem loop, as the n.40C is one of four bases stabilizing this loop and notably it pairs with n.46 G, a base previously reported to be mutated in MOPD1 (Kılıç et al., 2015). The other mutation, the 85-base-pair insertion in position n.101, is also predicted to have a major impact on conformation and to destroy the 3’ stem loop, and it is therefore also predicted to have a major functional impact on the snRNA [in-silico predictions made by Protein Data Bank 3SIU (New Jersey, USA) and PhyMol v.1.7 (Schrödinger, New York, USA) software].

**Discussion**

MOPD1 is generally described as a fatal condition within the first months or years of life. However, reports of less severely affected individuals are emerging (Abdel-Salam et al., 2012; Nagy et al., 2012).

Most of the previously reported cases were homozygous for the n.51 G>A mutation, a founder mutation in the Amish population, representing the most frequent genotype of MOPD1 patients and one associated with a shorter life span compared with other cases (Nagy et al., 2012). A patient described with a milder phenotype is compound heterozygous for n.66C>G and n.124 G>A mutations (Abdel-Salam et al., 2012). Thus, the fact that our cases present with a mild phenotype may be because of the compound heterozygous state or that one of the mutations lies outside the essential 5’ stem loop.

Seven of the previously reported mutations in patients with MOPD1 including n.30 G>A, n.46 G>A, n.50 G>A, n.50 G>C, n.51 G>A, n.53C>G and n.55 G>A mutations are located within the 5’ stem loop of the U4atac snRNA, a motif interacting with spliceosomal proteins. This secondary structure of the snRNA is highly conserved across species, suggesting that the site is of critical importance for minor U12 spliceosomal function (Edery et al., 2011). The 3’ stem loop is also believed to have an essential role, and complete deletion of the 3’ stem loop is reported to abolish the in-vivo splicing function of the minor spliceosome (Shukla et al., 2002).
The two mutations reported by us are both predicted by in-silico prediction tools to severely affect the secondary structure of the snRNA.

Our presented cases display some classical features of MOPD1, including prenatal and postnatal growth retardation, microcephaly, developmental delay, cataract, hearing loss and dysmorphic features, but unlike previously reported cases have survived into adult life.

Although no clear genotype–phenotype correlation exists on the limited number of milder cases reported previously and in our study, our findings at least indicate that the compound heterozygous genotype n.40C>T/n.101 tandem duplication (bp 16–100) in RNU4ATAC results in a mild phenotypic appearance of MOPD1 compatible with survival into adulthood. The presented cases further expand the mutational and phenotypic spectrum of the MOPD1 syndrome.

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Conflicts of interest

There are no conflicts of interest.