Association of Wolfram syndrome with Fallot tetralogy in a girl

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ABSTRACT

Wolfram syndrome (DIDMOAD: diabetes insipidus, diabetes mellitus, optic atrophy and deafness) is a rare neurodegenerative disorder. Mutations of the WFS1 (wolframin) on chromosome 4 are responsible for the clinical manifestations in majority of patients with Wolfram syndrome. Wolfram syndrome is also accompanied by neurologic and psychiatric disorders, urodynamic abnormalities, restricted joint motility, cardiovascular and gastrointestinal autonomic neuropathy, hypergonadotrophic hypogonadism in males and diabetic microvascular disorders. There are very limited data in the literature regarding cardiac malformations associated in children with Wolfram syndrome. A 5-year-old girl with Wolfram syndrome and tetralogy of Fallot is presented herein.

Key words: Wolfram syndrome, Fallot tetralogy, DIDMOAD.

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INTRODUCTION

First described in 1938 by Wolfram in four siblings, Wolfram syndrome (DIDMOAD) is a clinical condition involving concurrent diabetes insipidus, diabetes mellitus, optic atrophy and deafness. Usually, diabetes mellitus is diagnosed initially and followed by optic atrophy within the first 10 years of life and diabetes insipidus and optic atrophy during the second decade.

Mutations on the WFS1 gene on chromosome 4 are responsible for the clinical manifestations in majority of the patients with Wolfram syndrome. This gene encodes an endoplasmic reticulum membrane protein (wolframin) present in neurons, pancreatic β-cells, internal ear, heart, placenta, lung and liver. Although the function of this protein is unknown, its deficiency leads to increased stress within the endoplasmic reticulum, leading to impairment of cell cycle and calcium homeostasis. Wolframin has a role in maintaining the homeostasis of endoplasmic reticulum in pancreatic β-cells. Cardiac malformations in Wolfram syndrome have been reported rarely and may be responsible for the morbidity and mortality of the disorder. We present a case with Wolfram syndrome accompanied by tetralogy of Fallot and discuss the findings with reference to literature.

CASE

A 5.3-year-old female patient presented with excessive thirst and frequent urination with 3 kg of weight loss during the last month. She had been diagnosed with Fallot tetralogy at 4 months of age and had undergone surgery. In addition, audiometric examination had revealed bilateral sensorineural hearing loss when she was 3-years old and she was receiving oxybutynin treatment at 0.2 mg/kg/day for neurogenic bladder for the last 4 months. She was the second child of apparently healthy non-consanguineous parents, born at full-term by normal vaginal delivery following an uncomplicated pregnancy. Family history did not disclose diabetes mellitus.

Physical examination: weight: 14.9 kg (3-10th percentile), height: 122 cm (50-75th percentile), strabismus, and surgical scars on the thorax. Laboratory: plasma glucose was 314 mg/dl, urinary ketone negative, normal blood gases, serum C-peptide level 0.18 pmol/ml (0.15-1.10) and HbA1c value of as 11.2% The diagnosis was diabetes mellitus. Anti-insulin antibody was 4.5 U (0-8 U), anti-GAD 0.9 U/ml (<1 U/ml), and islet cell antibody negative. She was discharged under four doses of regular insulin (0.73 U/kg/day) treatment.

At the age of seven and a half, despite good glycemic control (most recent HbA1c, 6.92%), she developed excessive thirst, frequent urination, and nocturnal enuresis. Fluid intake was estimated to be 3500 cc/m²/day and urine output 9.4 cc/kg/h. Renal function tests (BUN: 8.9 mg/dl, creatinine: 0.37mg/dl) and serum
electrolytes (Na: 139 mmol/L, C: 5.1 mmol/L, Cl: 105 mmol/L) were normal. Serum osmolality was 275 mOsm/kg, urine osmolality 168 mOsm/kg, and serum ADH level <0.5 pmol/L. Central diabetes insipidus was considered and treatment with desmopressin nasal spray was started (10 µg bid). Desmopressin nasal spray therapy resulted in a marked improvement of excessive thirst, frequent urination, and nocturnal enuresis. Fundus oculi examination demonstrated bilateral optic atrophy and no sign of diabetic retinopathy. Pituitary magnetic resonance imaging did reveal absence of bright spot in the posterior pituitary. Wolfram syndrome was diagnosed accordingly.

Genomic DNA from proband and parents were extracted from peripheral blood leukocytes using the QIAamp DNA mini kit (Qiagen, 51304, Dusseldorf, Germany), according to standard procedures. Genomic fragments including coding regions and adjacent intronic regions of WFS1 were amplified with PCR, using previously described primers (genetic source). The amplicons were purified and analyzed with cycle sequencing with ABI BigDye Terminator Cycle Sequencing Kit v3.1 (ABI Applied Biosystems, Foster City, CA) on an automatic DNA sequencer (ABI 3130 Genetic Analyzer, Applied Biosystems). A known mutation (Y508fsX541, c.1523_1524delAT) was homozygous in exon 8 of WFS1 were detected in the proband (Figure 1). This mutation was present in heterozygous form in both parents.

**DISCUSSION**

Various genetic alterations are responsible for Wolfram syndrome. Majority of the cases have homozygous or compound heterozygote WFS1 mutations resulting in altered function of wolframin. However, recently, a dominantly inherited WFS1 mutation was found to underlie Wolfram syndrome in a Finnish family. Mutations in CISD2 have been identified in patients with a similar clinical picture but without diabetes insipidus (Wolfram syndrome 2). Similar to wolframin, the CISD2-encoded protein ERIS (endoplasmic reticulum intermembrane small protein) localizes to endoplasmic reticulum. Some patients with Wolfram syndrome demonstrate mutations at mitochondrial DNA. Mitochondrial defects leading to diabetes mellitus are usually accompanied by neurovascular disorders including mental retardation, migraine, deafness and convulsions. In our patient, Wolfram syndrome was associated with a homozygous mutation in WFS1.

Diabetes mellitus is generally the first clinical manifestation in Wolfram syndrome and develops during the first decade of life, at about 6 years of age in average. Since all of the features were not present, diagnosis of Wolfram syndrome could not be made in our patient at the onset of diabetes. Congenital rubella syndrome is known to be associated with type 1 diabetes, heart defects, sensory deafness, and cataracts.
Our patient had not such a history, cataract, or autoantibodies for type 1 diabetes. Recently, GATA6 mutations are reported to be a cause of severe congenital heart disease and neonatal/childhood-onset diabetes. Our patient neither experienced decreased exocrine pancreas function nor hepatobiliary abnormalities that occur with GATA6 mutations. Retrospectively, presence of congenital heart disease, sensorineural hearing loss in an autoantibody-negative child with diabetes and voiding dysfunction would have led to the diagnosis of Wolfram syndrome at the time of presentation since this rare association has been reported in relevant studies.

In summary, congenital heart defects can be a feature of Wolfram syndrome. This is particularly important for diagnostic approach of children with autoantibody-negative diabetes and incomplete features of Wolfram syndrome. Early diagnosis and appropriate management would prevent development of complications.

REFERENCES