INTRODUCTION

Oral-facial-digital syndromes (OFDs) are a group of heterogeneous syndromes characterized by abnormalities in the oral cavity, face and digits. To date, 13 types with different modes of inheritance have been distinguished based on characteristic clinical manifestations. The researchers reported a twelve-year-old male patient with the common features of oral, facial, and digital abnormalities of an OFDIX syndrome along with some unreported features. On assessment, it was found that he had a short stature and macrocephaly. The patient had sparse scalp hair and alopecia areata, which is reported only in females with OFDI. He showed hands postaxial polydactyly and unilateral bifid big toe. The patient displayed extraocular manifestations along with multiple intraocular findings. There were additional CNS findings in the form of subependymal and periventricular and arachnoid cysts. The arachnoid cysts were previously described in OFD I, II and III. This case overlapped with the clinical picture between OFD I, which is detected only in girls and OFD II.

CLINICAL REPORT

The patient for the present study is a twelve-year-old boy, the first born to consanguineous Somalian parents, with four sisters and 2 brothers. The family history is unremarkable. He was born at term following an uncomplicated pregnancy and delivery, to a healthy 20-year-old mother and a 27-year-old father, at the time of conception. The patient was referred to the Medical Genetics Clinic at King Abdulaziz University Hospital due to dysmorphism, delayed motor milestones and intellectual handicap. On assessment at the age of twelve years, he had a short stature (height was 113 cm at -5 SD), macrocephaly (skull circumference was 46 cm, >-3 SD) and was underweight (21.2 kg; <3th percentile). The patient had sparse hair with areas of alopecia (alopecia areata) (Fig. 1A), sparse eyebrows, transverse slanting palpebral fissures, ptosis, hypertelorism and apparent microphthalmia of the left eye (Fig. 1B). There was a broad nasal root and hypoplastic alar nasi with the columella extending below the nares. He had
Fig. 1. Clinical features.  A. Alopecia areata. B. Sparse eye brows, hypertelorism, ptosis.. C. Tongue nodules, bifid tip of tongue, and dental caries. D. Polydactyly of upper limb

Fig. 2. MRI brain. A&B: Multiple subependymal, periventricular and arachnoid tiny cysts. C&D: Left microphthalmia and retinal detachment, right retrobulbar focal cystic lesion and optic coloboma.
posteriorly rotated ears. The mouth was held open with a prominent lower lip and a high arched narrow palate, tongue nodules with a bifid tip, irregular hypertrophic gingiva and an alveolar ridge with dental caries, together with a history of operated tongue frenula (Fig. 1C). There was retromicrognathia. The hands and feet were short with brachydactyly and upper limb postaxial polydactyly (Fig. 1D), well formed in the right hand and rudimentary on the left side. The feet showed a broad first toe on the right side and bifid on the left side. The cardiac and chest examinations were normal. Genital examination showed hypospadius. The ophthalmological examination showed right side optic nerve coloboma and left microphthalmia, cataract, and total retinal detachment.

The chromosomal study reported a normal male (46, XY). The echocardiogram and the abdominal ultrasound were unremarkable. The complete blood count, coagulation profile, urea and electrolytes, thyroid function tests, liver enzymes, were all within normal limits. An X-ray of the feet revealed a small bony projection from the medial border of the left first toe (bifid toe) and medial inclination of bilateral metatarsals. An X-ray of the hands showed post-axial polydactyly. The brain magnetic resonance imaging (MRI) showed multiple subependymal, periventricular and arachnoid tiny cysts (Figs. 2A and B). The left eye showed microphthalmia and retinal detachment, while the right eye showed a retrobulbar focal cystic lesion and optic coloboma (Figs. 2C and D).

DISCUSSION

The OFDs from a pleiotropic effect of a morphogenetic impairment variably affecting the oral cavity, face, digits and other organs. To date, 13 types have been distinguished based on characteristic clinical manifestations, (Gurrieri et al. 2007; Adyanthaya and Adyanthaya 2015). OFD IX (MIM 258865) has been previously described by Gurrieri et al. (1992) who described two sibs as type II (autosomal recessive) with eye abnormality that overlaps type VIII, which share the same findings but has a different mode of inheritance (X-linked recessive) (Gurrieri et al. 2007). Since then there were seven additional reports of type IX (Jamieson and Collins 1993; Nevin et al. 1994; Sigaudy et al. 1996; Nagai et al. 1998; Tsai and O’Brien 1999; Erickson and Bodensteiner-er 2007). The patient comprised the common features of oral, facial and digital abnormalities of an OFD syndrome Gurrieri type (OFD IX) (Gurrieri et al. 1992) along with some unreported features. Known features are oral, facial, and digital findings, hypertelorism, tongue nodules with bifid tip, irregular hypertrophic gingival and alveolar ridge with dental caries. In this patient there was sparse scalp hair and alopecia areata, which is reported only in OFD I (Gurrieri et al. 2007; Chetty John et al. 2010). OFDIX (Gurrieri type) is normally characterized by paucity of digital findings; hallucal duplication was the only digital anomaly previously detected in this type, while this patient showed upper limb postaxial polydactyly which have not been previously reported in Gurrieri type. Also, Erickson and Bodensteiner (2007) reported the paucity of these findings in a series of cases (Gurrieri et al. 1992; Erickson and Bodensteiner 2007). In this study, the feet showed a broad big toe on the right side and bifid on the left side, which is a common finding in OFD IX.

Ophthalmological findings, which have been reported in OFD syndromes include hypertelorism, see-saw winking and strabismus. The most common intraocular finding described in OFDS IX is retinal coloboma as a distinctive feature (Gurrieri et al. 1992) followed by optic coloboma and a single case of retinal hamartoma (Tsai and O’Brien 1999; Erickson and Bodensteiner 2007). The patient in the present study displayed more than one finding in the form of left microphthalmia, cataract, and total retinal detachment. There were also right optic nerve coloboma and a retrolubular cyst. The presence of more than one ocular finding in one case has not been detected before.

CNS abnormalities previously detected in OFDIX included microcephaly, brain atrophy and dilated fourth ventricle (Gurrieri et al. 1992; Jamieson and Collins 1993; Sigaudy et al. 1996; Erickson and Bodensteiner 2007).

In this patient there were additional findings in the form of subependymal, periventricular and arachnoid tiny cysts. Arachnoid cysts were previously described in OFD II and III (Leao and Ribeiro-Silva 1995; Nagai et al. 1998; Gurrieri et al. 2007).

CONCLUSION

In conclusion this case showed an overlap of the clinical picture between OFD I and OFD
II. The patient showed oral abnormalities and alopecia areata which was similar to OFDI which is lethal in males as it is a X-linked dominant inheritance, however he had manifestations of OFDII in the form of polydactyly and CNS abnormalities, which are known to be less common in type IX. Also the patient had multiple eye abnormalities, which have been advocated as a specific finding of type IX. The presence of consanguinity and no findings in the parents may indicate an autosomal recessive mode of inheritance. This case describes the overlap with type of OFDII and may represent an allelic variant of OFD type IX or a new type of OFDs.

REFERENCES