First Report of 2q37 Microdeletion Syndrome in Morocco
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Abstract

2q37 microdeletion syndrome or Albright Hereditary Osteodystrophy like (AHO-like), also referred to as AHO3 or Monosomy 2q37qter, is characterized by developmental delay, intellectual disability, brachymetaphalangy of digits 3-5, short stature, obesity, hypotonia, characteristic facial appearance, autism, and skeletal defects. In this work, we report a case of cytogenetically visible de novo deletion of this 2q37 chromosome region, referred to genetic department of Mohammed VI University Hospital of Marrakech and we discuss its clinical features and laboratory tests. R-banding chromosomal analysis was performed for the patient from T lymphocyte cultures in vitro and blocked in metaphase. This technique showed a small terminal deletion of the long arm of chromosome 2 and the karyotype was designated 46, XX, del(2)(pter ---->q37:). Parental karyotypes (R-banding) were also realized, but didn’t show any chromosomal abnormality. So, in that case, genetic counseling is reassuring for the future pregnancies. We show through this work the interest of the geneticist in the diagnosis of rare syndromes and appropriate genetic counseling of patient and his family.

Keywords: 2q37 microdeletion; AHO-like; AHO3; monosomy 2q37qter; constitutional karyotype; genetic counseling; Mohammed VI University Hospital of Marrakech.

INTRODUCTION

2q37 microdeletion syndrome (or AHO-like, AHO3, Monosomy 2q37qter) is characterized by a round face, frontal bossing, flattened mid-face, deep-set eyes, epicanthic folds, anteverted nostrils, hypoplastic alae nasi, brachymetaphalangy of digits 3-5 (often digit 4 alone), minor ear anomalies, short stature and obesity. Hypotonia, and developmental delay are the most common manifestations of this syndrome [1]. Behavior problems are also frequent [2, 3] including hyperactivity, repetitive behavior, in many cases, these manifestations have been characterized as autistic disorders [4]. Other findings include seizures, congenital heart disease, Central Nervous System (CNS) abnormalities (hydrocephalus, dilated ventricles), umbilical/inguinal hernia, tracheomalacia, situs abnormalities, gastrointestinal abnormalities, and renal malformations.

In the present work, we report a case of 2q37 microdeletion syndrome or Albright Hereditary Osteodystrophy like syndrome, referred to Genetic department of Mohammed VI University Hospital of Marrakech and we discuss its clinical features and laboratory test results to correlate them with results found in the literature. In fact, this is the first report of 2q37 microdeletion syndrome in Morocco.

We show through this work the interest of the geneticist in the diagnosis of rare syndromes and appropriate genetic counseling of patient and his family.

CLINICAL CASE REPORT

The proposita aged one year, the last of two sibling (Fig.1). She was referred to genetic counseling of UTH-Mohammed VI of Marrakech by a pediatrician for facial dysmorphia associated with a delayed motor skills. Physical examination showed a height of 70 cm. Dysmorphology examination revealed a round face, a short nose with a depressed bridge, Mongoloid palpebral slit, hypertelorism, a front saillant, arched eyebrows with synophris, anteverted nostrils, Microstomia and thin lips with a long and a marked philtrum, retrongnathia, high-arched palate, hypoplasia of the right ear lobular (Fig.2). The examination of extremities showed brachydactyly of the hands (Fig.3). Radiographic examination of the cavum demonstrated
an imprint adenoid that projects on the posterior wall of the nasopharynx.

**Fig-1: Family pedigree**

**Fig-2: Facial dysmorphia**

**Fig-3: Picture of the hand**

**Fig-4: Karyotype of the patient (a), Combined karyotype of chromosome 2 (b)**

**CYTOGENETIC STUDY**

R-banding chromosomal analysis was performed for the patient from T lymphocyte cultures in vitro and blocked in metaphase. This technique showed a small terminal deletion of the long arm of chromosome 2 (Fig.4) and the karyotype was designated 46,XX,del(2)(pter----q37:). Parental karyotypes (R-banding) were also realized, but didn’t show any chromosomal abnormality.

**DISCUSSION**

Since 1983, several authors have described small terminal deletion of 2q associated with variable phenotypes [5- 9]. This deletion was visible on prometaphase karyotype in most cases published [10-13, 6, 14].

The clinical feature is characterized by severe facial dysmorphic which combines a round face, thinning hair, a prominent forehead, oblique palpebral fissures down and out, eyes deeply located in orbits, low supplied and arched eyebrows, midface hypoplasia, a depressed nasal bridge, low nose wings, advanced V-shaped nose, a thin upper lip and a high arched palate [1, 6]. In the present work, the patient present variable dysmorphic signs, brachydactyly, short stature and malformations. These criteria are already described in the clinical feature of the 2q37 deletion.

The dysmorphic features are even more numerous than the deletion is extended [1, 15] studied 5 patients with mental retardation and brachydactyly. He found that four had visible deletions and one had a submicroscopic deletion of 2q37.

Moreover, patients with 2q37 terminal deletion may have a small size in 23% and obesity. Malformations are also present (in 30% of patients) [16].

Generally, the variability of the clinical feature depends on the genes involved. According to the study
of [17], five genes are involved in AHO-like due to the deletion 2q37 especially; STK25, KIF1A and HDAC4. Moreover [18] are proposed HDAC4 as the gene in which a pathogenic variant caused the syndromic features of the 2q37 microdeletion syndrome.

The prevalence of the 2q37 microdeletion syndrome is unknown. This can be explained by the difficulty in recognizing the small terminal deletion on classical cytogenetic studies, and the failure to recognize the clinical syndrome on physical examination.

Diagnosis is based on cytogenetic analysis and molecular characterization. Research of translocation must also be performed, because the deletion may result from the transmission of a derivative chromosome.

The differential diagnosis includes other syndromes aneusomy segmental and Prader-Willi syndrome. The AHO (pseudo-hypoparathyroidism, PHP) and the pseudo-PHP (PPHP) should also be considered in the differential diagnosis but can be removed by highlighting of normal levels of calcium, phosphorus and parathyroid hormone in patients with the deletion 2q37 [18] demonstrated an overlap between 2q37 microdeletion and Smith-Magenis syndrome. Moreover [19] have found that two individuals with suspected phenotypic features of Kabuki syndrome had chromosomal rearrangements including a 2q37 deletion.

Clinical characteristics of 2q37 microdeletion syndrome are apparent and no cases of germinal mosaicism have been described nowadays. There appears a female predominance among patients with a 2q37 deletion [20], which is the case of our study.

2q37 microdeletion syndrome can be the result of a de novo chromosome abnormality or may be inherited from a parent who is a balanced translocation or inversion carrier [21] have shown that the majority of chromosomal terminal deletions were de novo (48/60 familial studies) and their parents have normal karyotype. In present work, no balanced translocation and no abnormalities were found in the karyotype of parents who want to have more children. In this case, genetic counseling is reassuring for the future pregnancies. However, when the karyotype of parents presents a balanced translocation, genetic counseling is not reassuring and prenatal diagnosis is possible for subsequent pregnancies. This diagnosis is possible by chromosome analysis of fetal cells obtained by amniocentesis usually performed at about 15 to 18 weeks’ gestation or by chorionic villus sampling (CVS) at about 10 to 12 weeks’ gestation. It is difficult to visualize terminal chromosome deletions in fetal cells, that's why we have to do the FISH technique to confirm the result.

The support of patient with 2q37 microdeletion syndrome should be multidisciplinary and include a full evaluation of major clinical signs. Speech therapy, physiotherapy and occupational therapy are necessary. Periodic reevaluation by a clinical geneticist to provide new recommendations and information about the syndrome is preferable. Moreover, the patient must perform periodic pediatric neurodevelopmental and behavioral assessments to help manage cognitive and behavioral problems.

CONCLUSION

Through this work, we highlight the contribution of the geneticist in the diagnostic approach and the identification of the terminal deletion 2q37 for appropriate support of patients and adequate genetic counseling for their families.

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REFERENCES