Case Report

3p partial trisomy and 13q partial monosomy with congenital malformations and psychomotor developmental delay

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ABSTRACT. We examined a girl presenting neuropsychomotor developmental delay and multiple malformations including antenatal and postnatal growth retardation, congenital heart defect, and facial dysmorphisms. Cytogenetic analysis was performed on peripheral blood lymphocytes with the GTG-banding technique, which revealed an unbalanced translocation: 46,XX,der(13)(13pter→13q34::3p24→3pter) pat. Karyotype analysis of the father demonstrated a balanced translocation, 46,XY,t(3;13)(p24;q34), indicating the inheritance of the derivative chromosome 13. The mother karyotype was normal. We suggest that most of the structural malformations seen in this patient are due to the 3p trisomy, while the neuropsychomotor alterations are a consequence of both chromosome aberrations.

Key words: Congenital malformations; Neuropsychomotor alterations; 3p trisomy; 13q monosomy
INTRODUCTION

Partial trisomy 3p is a rare chromosomal abnormality, and it can be either de novo or inherited from one parent with a balanced translocation. It has been known as a syndrome with multiple congenital abnormalities and intellectual deficiency, including neuropsychomotor retardation. Clinical manifestations are quite varied, depending on the amount of 3p material in the trisomic state. The partial 13q monosomy is a rare condition characterized by a wide range of clinical findings. Phenotype varies with the location and size of the deletion. It commonly produces malformations such as microcephaly, ear abnormalities, retrognathia, hypertelorism, palate fissures, hypotonia, and skeletal abnormalities, as well as psychomotor developmental delay and heart defects. We describe here a girl with both chromosomal alterations, and we suggest a correlation between each alteration and her phenotype.

CASE REPORT

We present a case report on a patient (LSS) of 19 months, who was the third child to a 23-year-old mother and a 45-year-old father, where the parents were unrelated. The mother had no familial history of congenital malformations or spontaneous miscarriage. The father had 20 other children and 3 of them had unknown causes of death.

The mother did not have any problems during the pregnancy, but she reported few fetal movements and amniotic fluid loss a few weeks before childbirth. She had a cesarean section, which was prolonged and dystocic due to the transverse position of the baby, who was cyanotic due to the intense suffering. Resuscitation was not necessary. The child had a 5-min Apgar score of 8 and normal newborn screening results. Birth weight was 3150 g.

The child had psychomotor developmental delay (speech and motor), microcephaly, large forehead, hypertelorism, strabismus, small nose with anteverted nares, depressed nasal bridge, prominent short philtrum, large mouth with downturned corners, prominent posteriorly rotated low-set ears, retrognathia, widely spaced inverted nipples, short sternum, right-convex thoracic scoliosis, and grade II systolic murmur (Figure 1). The patient was affected by dysphagia for any kind of food constitution, having a higher tolerance for liquids. Female genitalia was normal.

Figure 1. Clinical manifestations. A. Large forehead, small nose with anteverted nares, prominent short philtrum, large mouth with downturned angles, widely spaced nipples, and short sternum. B. Prominent low-set posteriorly rotated ears, retrognathia.
The patient had an acyanotic congenital heart disease (large ostium secundum-type interatrial communication, with moderate hemodynamic repercussion) and also a discrete muscular hypotrophy. She was affected by repetition bronchopneumonia since the 26th day of life, leading to multiple hospital admissions. She died in the 19th month of life due to respiratory problems.

High-resolution cytogenetic analysis was performed in the patient and her parents, after culture of the lymphocytes from their peripheral blood according to previous studies (Ford and Hamerton, 1956; Yunis, 1976). We studied 50 metaphases by GTG banding treatment. The mother’s karyotype was normal, while the father showed a balanced translocation (Figure 2), with a karyotype of 46,XY,t(3;13)(p24;q34). The patient inherited the father’s derivative chromosome 13, showing the karyotype 46,XX,der(13)(13pter→13q34::3p24→3pter)pat. The patient’s parents signed an informed consent approved by the Ethics Committee of Universidade Federal do Maranhão allowing the publication of the clinical data and photographs.

Figure 2. Father’s 3 and 13 chromosomes at band level 550.

Most of the clinical manifestations described in our patient are known to occur in both trisomy 3p and monosomy 13q syndromes, individually. Other phenotype manifestations are compatible with the clinical alterations related to 3p trisomy patients (Reiss et al., 1986; Chen et al., 2008; Ginocchio et al., 2008; Tan et al., 2011; Han et al., 2012), and some are related to 13q monosomy (Schinzel, 1983; Brewer et al., 1998; Brooks et al., 2006; Ballarati et al., 2007; Quélin et al., 2009), as shown in Tables 1 and 2.

The partial 13q monosomy commonly produces malformations such as microcephaly, ear abnormalities, retrognathia, hypertelorism, palate fissures, hypotonia, and skeletal abnormalities, as well as psychomotor development delay and heart defects (Table 1). The partial 3p trisomy has been known as a syndrome with multiple congenital abnormalities and intellectual deficiency, characterized by micrognathia, short neck, hypertelorism, large mouth with downturned angles, prominent philtrum, speech delay, congenital heart disease, and neuro-psychomotor retardation. Most patients die during the first two years of life (Table 2). Clinical manifestations are quite varied, depending on the amount of 3p material in the trisomic state.

In general, phenotypes associated with deletions are more severe than those associated with duplications, with a maximum tolerance in living organisms of 3 and 10% of the total genome, respectively (Brewer et al., 1998). Another study reported that patients with more distal deletions involving 13q33-13q34 do not show large structural malformations and growth deficiency (Schinzel, 1983). However, these patients are severely mentally handicapped. Therefore, it is most likely that the majority of the structural alterations manifested in our patient were caused by the partial 3p trisomy, while the psychomotor alterations were caused by both syndromes.
Finally, it should not be neglected that the father has a balanced translocation involving chromosomes 3 and 13, and even though he does not manifest any phenotypic abnormality, he has a higher probability of reproductive problems including miscarriages and descendants with chromosome abnormalities. The risk to carriers of balanced translocations has been shown to vary according to the chromosomal segments involved in the structural rearrangement. Therefore, genetic counseling for this family is highly recommended.

**Conflicts of interest**

The authors declare that they have no conflict of interest.

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