Ring Chromosome 13, A Rare Case Report

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Abstract

This case report describes the cytogenetic and molecular characterization of a child with de-novo ring chromosome 13 [r(13)]. The child presented with short stature, growth retardation and a Turner syndrome diagnosis. She was the first case of ring chromosome 13 cytogenetic alteration observed in our laboratory at Dicle University, Turkey. Her chromosomal composition was 46,XX,r(13). FISH (Fluorescence InSitu Hybridization) also confirmed the presence of r(13). A chromosomal microarray analysis using a CytoScan® Optima assay (Affymetrix) detected a 6.3 Mb deletion at 13q33.3q34. This rare case presented the first r(13) cytogenetic alteration detected in our laboratory.

Keywords: 13q deletion, Ring chromosome 13, karyotype, FISH, microarray

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Ring Chromosome 13, Nadir Bir Olgu Raporu

Öz
Bu olgu sunumunda de-novo ring kromozom 13 [r(13)] tespit edilen bir çocuk hastanın sitogenetik ve moleküler açıdan sonuçları değerlendirilmiştir. Hasta kısa boy, büyüme geriliği ve Turner sendromu ön tanısı ile laboratuvarımızda yönlendirilmiştir. Yapılan kromozom analizi sonucu 46, XX, r(13) kromozom kuruluşu edinilen hastaya FISH (Floresan InSitu Hibridizasyonu) analizi yapılarak r(13) varlığı teyit edilmiştir. CytoScan® Optima testi (Affymetrix) kullanılarak yapılan kromosomal mikrodizin analizi sonucunda 13q33.3q34'te 6.3 Mb'lik bir delesyon saptanmıştır. Bu nadir olgu, laboratuvarımızda saptanan ilk r(13) sitogenetik değişiklik olması nedeni ile önemlidir.

Anahtar kelimeler: 13q delesyonu, Ring kromozom 13, karyotip, FISH, mikrodizin.

INTRODUCTION

Deletions of chromosome regions are generally the result of double-stranded chromosome breaks, followed by the loss of the resulting acentric fragments during the next cell division. Such deletions can lead to ring chromosomes. A ring D chromosome was first described in a male subject in 1962 by Wang et al. There are considerable phenotypic variations in patients who carry a ring chromosome, whether complete or the mosaic form. Patients with ring chromosome 13 [r(13)] have various phenotypic abnormalities that correspond to specific breakpoints.

This case report presents the first instance of de-novo r(13) that has been analysed by our laboratory. We evaluated this rare case using cytogenetic, molecular cytogenetic and molecular techniques.

CASE PRESENTATION

An 11-month-old female child with short stature, growth retardation and a Turner syndrome diagnosis was referred to the Dicle University, Medical Faculty, Department of Medical Biology for karyotype analysis. The parents provided the child’s family history. According to the provided information, the subject was the second child of a couple who had been married for five years. There was no parental consanguinity, and the couple’s first child was a healthy three-year-old male.
Figure 2. Karyotype of the case.

The karyotypes of both parents were normal. A FISH analysis was performed using a whole chromosome staining probe (WCP), according to the Cytocell Standard protocol, and r(13) was confirmed (Figure 3).

Figure 3. FISH image of metaphase using Whole Chromosome Staining probe (WCP) and chromosome 13 was seen as green signal

A chromosomal microarray analysis using a CytoScan® Optima assay (Affymetrix) detected a 6.3 Mb deletion at 13q33.3q34. This deletion involved the following genes: LIG4 (601837), TNFSF13B (603969), IRS2 (600797), COL4A1 (120130), COL4A2 (120090), CASR2 (612800), ING1 (601566), ARHGEF7 (605477), SOX1 (602148), ATP11A (605868), MCF2L (609499), F7(613878), F10 (613872) PROZ (176895), PCID2 (613713), CUL4A (603137), LAMP1 (153330), ADPRHL1 (610620), TFDP1 (189902), ATP4B (137217), GRK1 (180381), GAS6 (600441), RASA3(605182), CDC16 (603461) and UPF3A (605350).

DISCUSSION

This report presents the first r(13) case detected by our laboratory in southeast Turkey. GTG banding of the metaphase chromosomes and the corresponding FISH analysis were performed. A terminal deletion was observed in the results of both methods; however, the banding karyotyping and FISH analysis were unable to define the precise breakpoint of the terminal deletion. Further investigation using chromosomal microarray analysis by CytoScan® Optima assay (Affymetrix) detected a 6.3 Mb deletion on the long arm of chromosome 13 (13q33.3q34).

Martin, Harvey and Pearn presented a study on r(13) syndrome that provided the detailed clinical and cytogenetic features of three unrelated cases. All these r(13) syndrome cases had breakpoints within the region bounded by bands 13q21 to 13q34 (5).

Brawn et al. reported that patients with interstitial deletions in the long arm of chromosome 13 had widely varying phenotypes. Based on cytogenetic analyses, they postulated that there was a specific region in 13q32 where deletions caused severe malformations, including digital and brain anomalies. To test this hypothesis at the molecular level, they studied chromosome 13 deletions in 17 patients; 5 had severe malformations, while there are 12 had only minor malformations. The results indicated that
the deletions in all the severely affected patients involved an overlapping region in q32, while the deletions in the mildly affected patients included some, but not all, of this overlapping region. They suggested that the severely malformed 13q- phenotypes resulted from deletions in a critical region of 13q32.

Liao et al. described a 10-month-old Chinese Han boy who presented with severe mental retardation, congenital bilateral hearing loss and multiple malformations. He had both r(13) syndrome and 47, XYY syndrome, and his karyotype was 47,XYY,r(13)(p11q34). After further research using high resolution, array-based, comparative genomic hybridization, a terminal deletion of 8.5 Mb was identified in the long arm of chromosome 13 (13q33.2—>q34). Liao et al. suggested that the patient’s hearing impairment might be a clinical feature associated with the distal 13q deletion and the r(13) formation. In addition, Ozsu et al. reported a case with ambiguous genitalia and r(13).

Minasi et al. described the first postnatal diagnosis of a child from Central Brazil with de novo cytogenetic alterations in 13q; this case was characterized by malformations of the brain, eyes, distal limbs and genitourinary tract, as well as a severe intellectual disability. The subject’s karyotype was a constitutive 46,XX,r(13)[77]/45,XX,-13[17]/46,XX,idic r(13), and a chromosomal microarray analysis detected a 15.39 Mb deletion. They suggested that further studies were needed to define whether genetic haploinsufficiency was associated with each major 13q deletion anomaly or whether one or more putative genes of the critical regions are contributing to these congenital malformations.

Ping-Chen et al. described prenatal diagnosis and molecular cytogenetic characterization of de-novo mosaic r (13) with fetoplacental chromosomal discrepancy.

**CONCLUSION**

Karyotyping is substantial to demonstrate autosomal chromosome anomalies but then the molecular methods should be used to define the exact break point of the terminal deletions.

**Congress presented:** This case reporte was presented on 26-29 October 2017 in Fethiye at National Medical Biology and Genetics Congress as seen below: S.Simsek, D.Oral, İ.Yücel, S.Tekeş, E.Unal, H.İsi.

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**Informed Voluntary Consent Form:** Consent form was taken from the patient.

**Declaration of conflicting Interest:** The authors declare that have no conflict of interest.

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