De Novo Three-Way Chromosome Translocation [46,XX,t(1;20;4)(p32;q12;q32)] in a Patient with Developmental Delay: A Case Report

Gelişme Geriliği Olan Bir Hastada Yeni Oluşum Üçlü Kromozom Translokasyonu [46,XX,t(1;20;4)(p32;q12;q32] : Bir Olgu Sunumu

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ABSTRACT

Complex chromosome rearrangements (CCRs) involve more than two breakpoints on two or more chromosomes are uncommon occurrences. We report a female with developmental delay with a complex three-way balanced translocation [46,XX,t(1;4;20)(p32;q32;q12)] identified by karyotyping. This paper is the first report of reciprocal translocation involving 1p, 4q and 20q associated with the developmental delay. The developmental delay may be due to the rearrangement of genetic material at these breakpoints and this incident may cause developmental delay.

Key Words: Chromosomal translocation, Developmental delay

ÖZET

Kompleks kromozom düzenlenmeleri ikisi ya da daha çok kromozomda ikiden fazla kırılma noktasını içeren sık olmayan oluşumlardır. Biz karyotip analizi sonucunda kompleks bir üçlü dengeli translokasyonu [46,XX,t(1;4;20)(p32;q32;q12)] gelişme geriliği olan bir kız çocuğunu rapor ediyoruz. Bu yazı 1p, 4q ve 20q içeren resiprokal translokasyon ile gelişme geriliğinin birlikte olduğu ilk raporudur. Sorun muhtemelen kritik bir ilişkiye sahip bu kırık noktalarında genetik materyalin yeniden düzenlenmesine bağlı olabilir ve bu olay gelişimsel gecikmeye neden olabilir.

Anahtar Sözcükler: Kromozomal translokasyon, Gelişim geriliği

INTRODUCTION

Reciprocal de novo translocations occur in about 1 in 2,000 newborns (1). Although balanced translocations are not often associated with abnormal phenotypes, unbalanced translocations resulting in deleted or altered gene sequences usually cause appreciable clinical features (2,3). Unbalanced translocations even more unusual are complex chromosome rearrangements (CCRs). CCRs are structural abnormalities that involve more than two break points and exchange of genetic material between two or more chromosomes (4,5). Almost all the carriers of the balanced reciprocal translocations are believed to be normal by phenotype. Moreover, it is known that the modification or inactivation of specific disease genes at chromosomal breakpoints have been very phenomenal in identifying genes that are associated with a variety of disorders, mostly early-onset disorders (6). The most common CCRs involve three chromosomes with breakpoints on each chromosome.

The occurrence of three-way multiple translocations is rare and often difficult to distinguish from balanced translocations without additional diagnostic tools such as fluorescence in situ hybridization (FISH) or microarray-based comparative genomic hybridization (array CGH) (7-9).

In the present paper, we report an 8-year-old female with developmental delay and myopathic face in whom we found a de novo complex translocation of 46,XX,t(1;4;20)(p32;q32;q12).

CASE REPORT

The proband is an 8-year-old female who was referred for developmental delay to our hospital. She was born at term by cesarean section to a 29-year-old mother. The proband's birth weight was 2200gr (<3rd percentile) and her length was 47cm (10th-25th percentile). The mother reports that she did not drink alcohol, smoke cigarettes, or use any unusual medications.
during the pregnancy. The father of the proband was 33-year-old with no prior family history of congenital abnormalities. At examination of the patient, she was 8-year-old, her weight was 16.4 kg (<3rd centile) and her height 112 cm (<3rd centile). An Auditory Brainstem Response test revealed bilateral normal sensorineural hearing. Cranial magnetic resonance imaging (MRI) revealed no pathology. Upon physical examination, the patient displayed minor dysmorphic features including simple ear, myopathic face, and clinodactyly (Figure 1A-C). Informed consent was received from parents for photographs. No other remarkable dysmorphic features or abnormalities were noted. Conventional cytogenetic analysis was performed on stimulated 72-hour culture of a peripheral blood specimen. The cells were cultured and processed by conventional methods and the chromosomes were stained with Giemsa–Trypsin–Giemsa banding (GTG). The karyotype was determined to be 46,XX,t(1;4;20)(p32;q32;q12) (Figure 2). The parents had normal chromosomal structure. Similarly, the first female child showed normal 46,XX karyotype. The family history was unremarkable.

**DISCUSSION**

The common cause of mental retardation and the wide range of physical abnormalities is balanced chromosome rearrangement. As such, it is difficult to interpret, posing as a diagnostic challenge in human development. We report the cytogenetic finding of a patient with a complex chromosome rearrangement involving 1p32, 4q32 and 20q12 showing an association with developmental delay, simple ears, and clinodactyly. This association has not been reported previously. Failure to detect the correct chromosomal recipient would have resulted in false results of the prenatal diagnosis performed in the second and third pregnancy of the consultand. This finding stresses the importance of cryptic terminal translocations in clinical cytogenetics. Balanced chromosomal translocations may cause damage or alteration of the functional genes at the breakpoints of the defective chromosomes resulting in the disease phenotype (10). It was described previously that children who inherit reciprocal balanced translocation from one of the parents show association with congenital malformation (11-13).
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The risk for offspring of a carrier of a balanced CCR is difficult to estimate (14). Batista et al gave a risk of miscarriage of about 50% based on an extended review of 30 families with CCRs (15). Gorski et al. analysed 25 CCR families with 67 informative pregnancies and found a risk of abnormalities of 18.4% for live births (16). Nearly 50% of all liveborn offspring were also CCR carriers. The overall risk of an abnormal outcome of pregnancy seems to be about 50%.

In carriers of balanced translocation, the possible reason for the association of congenital malformations could be gene inactivation or disruption at the breakpoint or a position effect (17). However, in the case of inherited reciprocal translocation seen in the proband, the breakpoint could inactive genes, subsequently unmasking a recessive allele inherited from the other parent (18). The other possible reason could be the occurrence of unequal crossing-over during meiosis that may have resulted in submicroscopic duplications or deletions, as proposed by Jacobs (17).

The deletion may be causing a position effect on the chromosome and may play a role in the proband’s phenotype, either because the chromosomal rearrangement separated a promoter from its transcriptional regulatory element, resulting in gene silencing; the rearrangement juxtaposed a gene with an enhancer from another gene, leading to inappropriate gene expression; or the rearrangement moved a gene and its regulatory elements to a region of the genome that is transcriptionally silent, such as heterochromatin. Alternatively, the translocation breakpoints may have interrupted a gene or genes.

In view of an increased risk of having congenitally abnormal children, carriers of balanced reciprocal translocation should therefore be advised to seek genetic counseling. The genetic counseling for a balanced translocation carrier is often difficult and may require some caution, especially when the fetal karyotype is balanced.

REFERENCES


