THE EFFECT OF FAMILIAL BALANCED RECIPROCAL TRANLOCATION t(9;11)(p12;p11.2) TO REPRODUCTIVE PERFORMANCE

Gülser ÖKTEM\textsuperscript{1}, Nurten KARA\textsuperscript{1}, Sengül TURAL\textsuperscript{1}, Davut GUVEN\textsuperscript{2}, Nevin KARAKUS\textsuperscript{1}

\textsuperscript{1} Department of Medical Biology and Medical Genetics, Ondokuz Mayis University, Faculty of Medicine, Samsun, Turkey

\textsuperscript{2} Department of Gynecology and Obstetric, Ondokuz Mayis University, Faculty of Medicine, Samsun, Turkey

SUMMARY

Aim: In this study, we report a couple who had been infertility problem for eight years and they had four failed IUI and one failed IVF. In second IVF attempt the women got pregnant and it resulted with abortion in eight week.

Material and methods: Cytogenetic analysis was performed by standart peripheral blood culture and GTG method by using phytohemagglutinin-stimulated lymphocyte.

Results: The women and her husband were phenotypically normal but karyotype analysis revealed 46, XX and 46,XY,t(9;11)(p12;p11.2) respectively. The mother of the husband's karyotype analysis showed the same translocation. The father of the husband's could not examined because he was not alive. Also two uncles of the husband were suffered from infertility for fifteen and five years respectively. However cytogenetic analyses of the uncles had not been accomplished yet. Balanced translocation carriers could give unbalanced chromosomes.

Conclusion: The infertility cases of especially monosomy 9p and the other similar translocations that can result from the parents of balanced translocation carriers are represented here by comparing.

Key words: chromosomal translocation, infertility


AİLESEL DENGELİ RESİPROKAL TRANSLOKASYON t(9;11)(p12;p11.2)'UN ÜREME PERFORMANSINA ETKİSİ

ÖZET

Amaç: Bu çalışmadada, sekiz yıldır infertil, dörtlü IUI (Intrauterin Inseminasyon) denemesinin başarısız, IVF (In Vitro Fertilizasyon) denemesinin icinicide başarılı olan fakat 8 haftalıkken gebelik kaybı yaşayan bir çift sunuldu.

Gereç ve yöntemler: Sitogenetik analizde, periferik kandan elde edilen kromozomlara tripsin gizma bantlama (GTG) uygulanarak karyotip analizleri yapıldı.

Bulgular: Fenotipik olarak normal görülen olguların karyotipleri 46,XX ve 46,XY,t(9;11)(p12;p11.2) olarak saptandı. Erkek olgunun annesinde de aynı translokasyon bulundu. Baba hayatta olmadığından incelenemedi. Ayrıca olgunun iki daysının oglarından biri 15 yıllık evli ve infertil, diğer 5 yaşlı evli ve infertildir, bu kişilerin hıza ulaştılamadığından incelenemedi.

Tartışma: Dengeli translokasyonu olgumuz verebileceğini gametlerden özellikle 9p monozomisi ve infertilitenin birlikte görüldüğü diğer benzer translokasyonu olgular karşılaştıralar sunulmuştur.

Anahtar kelimeler: kromozomal translokasyon, infertilite


Address for Correspondence: Dr. Gülser Öktenc. Ondokuz Mayis Üniversitesi Tıp Fakültesi, Tibbi Biyoloji Anabilim Dalı, 55139 Samsun, Turkey

Phone: + 90 (533) 492 52 82

e-mail: stural@omu.edu.tr

Received: 04 August 2011, revised: 30 November 2011, accepted: 11 January 2012, online publication: 12 January 2012

DOI: 10.5505/tjod.2012.14227
INTRODUCTION

Several studies have been shown that, aneuploidy, translocations, inversions, deletions of the Y chromosome and DNA damage may be effective in infertility(1). Chromosomal anomaly frequencies reported in the general population are lower than 1%, whereas patients with reproductive problems groups in around 5%. Among these changes, autosomal Robertsonian translocations and chromosome aberrations are ranks first in(2). Chromosomal anomalies are more common in infertile men although the cause of infertility in both men and women can(3). Especially in infertile male patients, incidence of somatic chromosomal abnormalities have been reported between 2.2% and 19.6. Sex chromosome abnormalities are the most common in human chromosomal abnormalities. The rate of sex chromosome abnormality in infertile men is 3.8%, while the abnormality rate of autosomal chromosomes are 1.3%(4). When infertile males compared to those of normal, the incidence of structural chromosomal abnormalities in men 1-4 times higher(1-5). These anomalies influence the reproduction. If the number of sperm reduces, incidence of anomaly increases. In oligospermia group, the most common anomalies are autosomal anomalies (3%), while in azoospermia group sex chromosome abnormalities (12.6%) are dominant(6). Changes in autosomal chromosomes, especially in Robertson-type translocations, during spermatogenesis may disturb some central effects of sperm density. Similar anomalies in the female carriers gametogenesis appears, however, is not affected. Therefore, in female carriers there is a risk of having spontaneous abortion or malformed children. However, male carriers of chromosomal abnormalities may cause sterility or subfertility(7). In this study, a family carrier of chromosomal t(9, 11)(p12: p11.2) with reproductive problems were presented.

CASE REPORT

Informed consent was taken all from family members and conventional cytogenetic methods and GTG (Giemsa-Trypsin) banding techniques were applied. Chromosome analysis were done, 30 metaphases were examined and 10 metaphases were karyotyped. Karyotypes were described according to the International System for Cytogenetic Nomenclature 2009(8). We presented a couple who have been suffered from infertility for eight years. After 4 failed IUI (IntrauterineInsemination), the second IVF (In Vitro Fertilization) was successful but it resulted 8 weeks of pregnancy loss. The couple karyotype results were 46,XX and 46,XY, t(9;11)(p12;p11.2). In family examination, mother of the male case who has the same translocation, could got pregnant after three years and first one was stillbirth and second one was first trimester pregnancy loss. After these, she had got a healthy child (Generation III, individual 11). Father of male case could not examined because he is not survive. Also generation III individual 2 married for 15 years and generation III individual 6 married for 5 years were also suffer from infertility but we could not reached them and we could not examine their karyotypes (Figure 2).

Table 1: Karyotype of the case.

<table>
<thead>
<tr>
<th>Table 1: Karyotype of the case.</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Chromosome</strong></td>
</tr>
<tr>
<td><strong>Individual</strong></td>
</tr>
<tr>
<td><strong>Karyotype</strong></td>
</tr>
<tr>
<td><strong>Generation I</strong></td>
</tr>
<tr>
<td><strong>Generation II</strong></td>
</tr>
<tr>
<td><strong>Generation III</strong></td>
</tr>
<tr>
<td><strong>Generation IV</strong></td>
</tr>
</tbody>
</table>

Table 2: Pedigree of the family.
DISCUSSION

Translocations are more common in infertile males than normal males\(^9\). These kinds of chromosomal rearrangements cause formation of unbalanced gametes by negatively affecting the spermatogenesis. These unbalanced gametes may cause disorders like recurrent pregnancy lose, congenital malformation, delayed development and mental retardation.

In a 46,XX,t(9;11)(p22;p15.5) case with different breakpoints from our case, the SRY gene was assigned to be positive, and delayed development and sex reversal were also observed\(^{10}\). Further, an infertile case with a 46,XY,t(9;15)(p10;q10) karyotype was defined\(^{11}\). Gonadal disgenesis and sex reversal (phenotypically male) were also observed in a case with 9p monosomy. Cases of infertility and sex reversal with different translocations of chromosome 9 were reported; t(9;13), t(9;3), t(7;9), t(4;9), del9(p23)\(^{12-17}\). In various leukemia patients, t(9;11) cases with different break points were observed\(^{18,19}\). Infertility cases with balanced translocations of different chromosomes were also reported\(^{4,6,11,20}\). The common results of these findings were the possible association of chromosomal abnormalities with incorrect chromosome coupling and crossing over in meiosis. The other possibility is the prohibition of the genes, related to testicular development and function, in chromosomal break points. The translocation in our case did not influence spermatogenesis, because the result of spermiogram test was normal. TET1 (testis-expressed transcript) gene was mapped in the 9p21-22 band region which was in the break point of our case\(^6\). The translocation of TET1 to chromosome 11 might cause infertility by showing a different position effect. As a result, in the translocated region, a detailed molecular analysis is needed to be done because of the possible cause of this region to infertility, especially in males.

Conclusion

In this study, it was emphasized that, chromosomal translocations might have an important role in the etiology of infertility and a detailed molecular analyses of genes, inside the related break points, had to be done.

KAYNAKLAR

15. Hoo, J.J., Salafsky, I.S., Lin, C.C. Possible localisation of a


