Distinct mechanism of formation of the 48, XXYY karyotype

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Abstract

Background: To expose the unusual nature of a coincident sex chromosomal aneuploidy in a patient and his father. Molecular mechanisms involved probably are based on the sperm chromosome of paternal origin, which determine the mode of formation. Conventional cytogenetics techniques and multiple Quantitative Fluorescent PCR of STR markers in sexual chromosomes in the patient and his parents.

Results: 48,XXYY and 47,XXY aneuploidies in the patient and his father, respectively, were identified. The additional X and Y chromosomes showed parental origin.

Conclusions: An infrequent origin of the 48,XXYY syndrome was demonstrated. Mostly, it is thought to result from an aneuploid sperm produced through two consecutive non disjunction events in both meiosis I and II in a chromosomally normal father, but in our father’s patient a 47,XXY was discovered. It is suggested that a higher incidence of 24,XY and 24,YY sperm may be possible in 47,XXY individuals and an increased risk for aneuploidy pregnancies may exist. Although 48,XXYY patients and Klinefelter syndrome are often compared, recently they are regarded as a distinct genetic and clinical entity.

Keywords: 48,XXYY, 47,XXY, Mechanism origin, Paternal, Spermatogenesis

Background

The 48,XXYY Syndrome is an uncommon sex chromosome aneuploidy condition that is characterized by the presence of one extra X and Y chromosomes in males. Until recently, XXYY Syndrome was considered as a variant of the Klinefelter syndrome (which includes the karyotypes 47,XXY; 48,XXYY; 48,XXXY and 49 XXXXY), but numerous authors have become aware that it has distinctive features. 48,XXYY syndrome the first observation was in 1960 by Muldal and Ockey [1] and later by Court Brown et al. [2] was described in 1964 and occurs in an estimated 1:18,000–1:50,000 males [3]. The patients show similar features of both individuals carrying 47,XXY and 47,XXY karyotypes [4], including tall stature, narrow shoulders, broad hips, sparse body hair, gynecomastia, normal to slightly decreased verbal intelligence [4,5] and hypergonadotrophypogonadism and infertility based on: small testes, absent spermatogenesis, normal to moderately reduced Leydig cell function, increased secretion of follicle-stimulating hormone and androgen deficiency [4-6].

When 48,XXYY and 47,XXY patients are clinically compared some differences are found: The first condition shows more frequently facial dysmorphism and congenital malformations [7] and also have a lower IQ with frequent and severe behavioral and psychiatric problems, including attention deficit hyperactivity disorder, autism spectrum disorders, and mood and tic disorders [6,7]. Males with sex chromosomal aneuploidies are known to have variability in their developmental profile with the majority presenting with expressive language deficits [6-8].

During the formation of the sperm, aneuploidy of the sex chromosomes is more frequent than aneuploidy of any of the autosomes and an association has been established between increased sperm aneuploidy and male subfertility [9].
A direct relationship between the number of additional sex chromosomes and the severity of the phenotype is generally assumed [10].

Some studies suggest that the frequency of XY sperm increases with age in parents of children with Klinefelter syndrome [11]. If one of these atypical sperm contributes to the genetic makeup of the child, it will have an extra X and Y chromosome in every cell.

47,XXY patients have a prevalence of approximately 0.1% of male population with the extra Y chromosome resulting from paternal non-disjunction at meiosis II. Usually they are fertile, with most of sperm cells with a normal karyotype. It has been hypothesized that one of the two Y chromosomes is lost before entering spermatogonia in meiosis. Several sex chromosome pairing configuration have been described at the pachytene stage: X + YY (most frequent), XX + Y, X + Y + Y and XYYtrivalent [12-14]. Only XYY trivalent configuration is believed to produce sex chromosome aneuploid sperm. An inverse correlation has been found between the proportion of XYY viable germ cells and low sperm counts with elevated sperm sex aneuploidies. Sex chromosome aneuploidies are more frequently of paternal origin as follow: 50% of 47, XXY, 100% of 47,XXY and 70-80% of 45,X [15].

Most of 48,XXXY are thought to result from an aneuploid sperm produced through two consecutive non-disjunction events in both meiosis I and II in a chromosomally normal father, so a boy with XXXY has one X chromosome from his mother and the additional XYY from his father.

We reported a case of a child with karyotype 48,XXYY, whose neurodevelopment delay, poor motor coordination, behavioral problem, facial features and unilateral kidney was the indication for genetic testing at 7 years of age. One case of 48,XXXY syndrome with unilateral renal aplasia has been previously reported [16].

Father's karyotype was determined and surprisingly, the result was 47,XXY. This is the first case of a 48, XXXY patient whose father carries a sex aneuploidy. Parental origin and hypothesis about the origin of the additional X and Y chromosomes in the patient were performed based on molecular data.

**Results**

The QF-PCR results confirmed the obtained data extracted by using the conventional cytogenetics techniques. The abnormal karyotypes with additional X and/or Y chromosomes: the 48,XXXY and 47,XYY karyotypes were found in all metaphases of the patient and his father, respectively (Figure 1).

The mother had a normal karyotype 46,XX and also the patient's brother showed anormal 46,XY karyotype. Size in base pairs and ratios for the area corresponding to the peaks of the fluorescence signal detected X-chromosome markers amplified specific STR, demonstrated that the patient had received one X chromosome from the mother and the other from the father. The extension of the molecular characterization (Table 1) of the 48,XXYY karyotype to his parents revealed the origin is paternal, as is generally accepted in this syndrome. The Figure 2 shows the marker image X22 (2:1:1 patient) and (2:1 patient’s father) and SRY gene amplification. In the (QF-PCR) showed an envelope in 2X/2Y patient and 1X/2Y his father. This results in a sperm with three sex chromosomes (one X chromosome and two Y chromosomes).

We propose two possible origins of the extra sex chromosomes in the patient (spermatogonia with a normal karyotype or with an extra Y chromosome).

**Discussion and conclusions**

Molecular and cytogenetic characterization of the patient carrying the 48,XXXY karyotype indicated a paternal origin of the additional X and Y chromosomes. Although there appears to be no particularly increased risk that a 47,XXY male will have a chromosomally abnormal child, it is not possible to discard the effect of the paternal aneuploidy in this patient.

It is accepted that XYY spermatogonia are able to produce altered 24,XY and 24,YY spermatozoa, although the ejaculate of XY men have generally shown a very small increased fraction of them. In a small percentage of patients, mitotic non-disjunction occurring during the early stages of a 46, XY zygote can result in 48,XXYY syndrome [17].

We propose two possible origins of the extra chromosomes in 48,XXYY patient: it is accepted that extra chromosome Y is lost in most of primary spermatocytes (pachytene cells) of 47,XXY patients. Also, it has been found that a high proportion of XYY configuration at pachytene in testis is negatively correlated with the sperm count in ejaculates [18,19], so the minority of oligospermic patients has a higher incidence of disomic 24,XY and 24,YY spermatoocytes.

In our patient's father there is no history of infertility so we can think that the extra Y chromosome is lost in most of spermatocytes. In those chromosomally normal sperm precursors (46,XY cells), two consecutive nondisjunction events in both meiosis I and II could originate an abnormal 25,XY sperm. This option also discards the implication of the karyotype character of the father in the appearance of sex chromosome abnormality in the child.

As we have commented previously, several sex chromosome pairing configurations have been described at the pachytene stage, and only XYY trivalent configuration is believed to produce sex chromosome aneuploidy. Spermatocytes with a trivalent configuration are compatible with the progress of meiosis [20,21]. Other sex chromosome configurations are not observed in Spermatocytes I with apoptosis occurring at the testicular level [19].
the case of XYY configuration, we can speculate that a single non disjunction event of all gonosomes in meiosis I of these cells could also originate a 25,XYY sperm. If this has been the mechanism involved in our patient, an increased risk for aneuploidy pregnancies may exist.

The QF-PCR technique resulted extremely sensitive to detect X and Y chromosome anomalies in postnatal diagnosis. The case presented by Zhang and Li (2009) [22] also shows that QF-PCR assays can provide rapid prenatal diagnosis of numerical sex chromosome aneuploidy.

It has been suggested that additional molecular data gained increased importance in this variant of Klinefelter syndrome, in which distinct patterns of X inactivation could play a role in the observed differences in the degree of clinical manifestations of patients. The wide range of psychological and medical problems has been attributed to skewed X chromosome inactivation [23].

Metabolic syndrome is associated with insulin resistance, the leading cause of truncal obesity and muscle

Table 1 Molecular and cytogenetic characterization

|                | Patient          | Father          | Mother         |
|----------------|------------------|-----------------|----------------|
| KARYOTYPE      | 48,XXYY          | XYY (QF-PCR)    | XX (QF-PCR)    |
| AMEL (Xp22.31-| XY               | XY              | X              |
| Xp22.1/Yp11.2) |                  |                 |                |
| SRY (Yp11.31) | Present          | Present         | No             |
| X22 (Xq28Yq)  | 199/225 bp       | 199/225 bp      | 220/241 bp     |
|               | 241 bp(2:1:1)    | bp(2:1)         |                |
| DXS6854 (Xq26.1) | 110/110 bp      | 110 bp          | 106/110 bp     |
|               |                  |                 |                |
| XHPR (X26.1)  | 274/285 bp       | 285 bp          | 274/281 bp     |
|               |                  |                 |                |
| DXS6803 (Xq21.31)| 110/120 bp      | 110 bp          | 120/120 bp     |
|               |                  |                 |                |
| DXS8377 (Xq28) | 250/262 bp       | 250 bp          | 253/262 bp     |
|               |                  |                 |                |
| DXS6809 (Xq21.33) | 263/271 bp     | 263 bp          | 267/271 bp     |
|               |                  |                 |                |
| G10_STS47 (Yq11.22) | Present  | Present         | No             |

Set of DNA fragment containing genes and genetic markers Short Tandem Repeat (STR) analysed, using Quantitative Fluorescence PCR in the patient and his parents. Parental alleles of the polymorphic markers evidenced that the additional X chromosome of the patient came from the father.
wasting, which was observed in the described patient. Also considered an endocrine disorder [24,25] and the risk of developing type 2 diabetes is high.

48,XXYY resembles Klinefelter’s syndrome although they are now considered different entities. It has been suggested that parent of origin of extra chromosome in 47,XXY influences the phenotype. Additional studies in 48,XXYY patients are needed to elucidate any epigenetic factors affecting the phenotype. Increased awareness of the developmental, psychological, and medical features of 47,XXY and 48,XXYY is important to facilitate timely diagnosis and initiation of appropriate screenings and treatments that are important for optimal outcomes [4,8,26].

Material and methods

Patient
The patient was a Caucasian Spanish 7 year old, who was referred to the Genetics Unit of the Infanta Cristina Hospital in Badajoz. This patient was referred for karyotyping due to hypergonadotrophic hypogonadism, tall stature, and other associated features include neurodevelopmental delay, poor motor coordination, behavioral problem, facial features and unilateral kidney, specifically, anorchia with unilateral renal agenesis. Right orchidopexy intervene exeresis of the left cord.

The patient has a long face, epicanthal folds, hypertelorism, narrow palpebral fissures, full lips and dental defects. Overweight was evident at 5 years of age and obesity from the 7 (3.6 standard deviation in weight), parents are not obese. The psychomotor and neurocognitive developmental were normal. No deviations were observed in progression in the areas of communication and socialization, language and motor adaptive wide. For all this clinically suspected to have Klinefelter syndrome.

Methods

After obtaining informed consent of the parents’ proband, peripheral blood was extracted and processed by standard techniques for conventional cytogenetic and molecular analysis. Quantitative fluorescent PCR (QF-PCR) assay were carried out in the patient, her parents and his unique brother. Chromosomal analysis was performed using standard techniques. Giemsa-based chromosomal banding and staining techniques detected the numerical chromosomal abnormalities.

Genomic DNA was obtained from the patient, parents and brother. Multiple quantitative fluorescent PCR (QF-PCR) was used including the amplification of amelogenin (AMELX and AMELY), which is presented on both sex chromosomes in a biallelic form, SRY gene (sex-determining region Y), a polymorphic short tandem repeat (STR) on the pseudoautosomal region of X and Y (X22), five polymorphic X-specific STRs (DXS6803, XHPR, DXS6803, DXS8377 and DXS6809), and a Y-specific marker (G10_STS47). STRs result in fragment length and area ratio differences which were analyzed with Applied Biosystems 3130 Genetic Analyzer and GeneMapper® Software Version 4.0. GeneMapper performed the identification of DNA fragments of interest and calculated their size in nucleotides, height and peak area. The size alleles corresponding to these peaks were identified and the results obtained in the child were compared and attributed to paternal or maternal ones for each marker; the ratio of heights or peak areas concerned.

Competing interests
The authors declare that they have no competing interests.

Authors’ contributions
RRL and JMC conceived the study and participated in its design and coordination. Helped to draft the manuscript. AMB carried out the molecular genetic studies, participated and drafted the manuscript. MGC, THM and EBB participated in the study design and protocol development. All authors read and approved the final manuscript.

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References
1. Muldal S, Ockey CH, Thompson M, White LL: ‘Double male’-a new chromosome constitution in the Klinefelter syndrome. Acta Endocrinol (Copenh) 1962, 39:183–203.
2. Court Brown WM, Jacobs PA, Price WR: Sex chromosome aneuploidy and criminal behaviour. Eugen Soc Symp 1968, 4:180–193.
3. Sorensen K, Nielsen J, Jacobsen P, Ralle T: The 48 XXXY syndrome. J Ment Defic Res 1978, 22(4):197–205.
4. Viscosatto J, Graham JM Jr: Klinefelter syndrome and other sex chromosomal aneuploidies. Orphanet J Rare Dis 2006, 2013, 1:1–42.
5. Frühmesser A, Kotzot D: Chromosomal variants in Klinefelter syndrome. Sex Dev 2011, 5(3):109–123.
6. Tartaglia N, Davis S, Hench A, Nimishakavi S, Beauregard R, Reynolds A, Fenton L, Albrecht L, Ross J, Viscosatto J, Hansen R, Hagerman R: A new look at XXXY syndrome: medical and psychological features. Am J Med Genet A 2008, 151(2):1509–1522. 146A.
7. Tartaglia N, Ayari N, Howell S, D’Epagnier C, Zeitler P: 48, XXXY and 49, XXXXY syndromes: not just variants of Klinefelter syndrome. Acta Paediatr 2011, 100(6):851–860.
8. Viscosatto J, Graham JM Jr: Social function in multiple X and Y chromosome disorders: XXX, XXXY, XXXXY. Dev Disabil Res Rev 2009, 15(4):328–332.
9. Mosani N, Pattinson HA, Carter MD, Cox DM, Raddermark AW, Martin RH: Chromosomal analysis of sperm from men with idiopathic infertility using sperm karyotyping and fluorescence in situ hybridization. Fertil Steril 1995, 64(4):811–817.
10. Linden MG, Bender BG, Robinson A: Sex chromosome tetrasomy and pentasomy. Pediatrics 1995, 96(4 Pt 1):672–682.
11. Lowe X, Eskenazi B, Nelson DO, Kidd S, Alme A, Wyrobek AJ: Frequency of XY sperm increases with age in fathers of boys with Klinefelter syndrome. Am J Hum Genet 2001, 69(5):1046–1054.

12. Mercier S, Morel F, Roux C, Clavequin MC, Bresson JL: Analysis of the sex chromosomal equipment in spermatozoa of a 47, XYY male using two-colour fluorescence in-situ hybridization. Mol Hum Reprod 1996, 2(7):485–488.

13. Rodrigo L, Peinado V, Mateu E, Remohí J, Pellicer A, Simón C, Gil-Salom M, Rubio C: Impact of different patterns of sperm chromosomal abnormalities on the chromosomal constitution of preimplantation embryos. Fertil Steril 2010, 94(4):1380–1386.

14. Chevet E, Rousseaux S, Monteil M, Usson Y, Cozzi J, Pelletier R, Selé B: Meiotic behaviour of sex chromosomes investigated by three-colour FISH on 35,142 sperm nuclei from two 47 XYY males. Hum Genet 1997, 99(3):407–412.

15. Templado C, Vidal F, Estop A: Aneuploidy in human spermatozoa. Cytogenet Genome Res 2011, 133(2-4):91–99.

16. Zantour B, Sfar MH, Younes S, Alaya W, Kamoun M, Mkaouar E, Jerbi S: 48XXYY Syndrome in an adult with type 2 diabetes mellitus, unilateral renal aplasia, and pigmentary retinitis. Case Report Med 2010. doi:10.1155/2010/612315.

17. Katulanda P, Rajapakse JR, Kariyawasam J, Jayasekara R, Dissanayake VH: An adolescent with 48,XXYY syndrome with hypergonadotrophic hypogonadism, attention deficit hyperactive disorder and renal malformations. Indian J Endocrinol Metab 2012, 16(5):824–826.

18. Solari AJ, Rey VG: The prevalence of a YY synaptonemal complex over XY synapsis in an XYY man with exclusive XYY spermatocytes. Chromosome Res 1997 Nov, 5(7):467–474.

19. Milazzo JP, Rives N, Mousset-Siméon N, Macé B: Chromosome constitution and apoptosis of immature germ cells present in sperm of two 47, XYY infertile males. Hum Reprod 2006, 21(7):1749–1758.

20. Blanco J, Egizazu J, Vidal F: Meiotic behaviour of the sex chromosomes in three patients with sex chromosome anomalies (47, XXX, mosaic 46, XX/47, XXX and 47, XXY) assessed by fluorescence in-situ hybridization. Hum Reprod 2001 May, 16(5):887–892.

21. Rives N, Siméon N, Milazzo JP, Barthélémy C, Macé B: Meiotic segregation of sex chromosomes in mosaic and non-mosaic XYY males: case reports and review of the literature. Int J Androl 2003 Aug, 26(4):242–249.

22. Zhang QS, Li DZ: A case of 48, XXXY syndrome detected prenatally by QF-PCR. J Matern Fetal Neonatal Med 2009, 22(12):1214–1216.

23. Itsuka Y, Bock A, Nguyen DD, Samango-Sprouse CA, Simpson JL, Bischoff FZ: Evidence of skewed X-chromosome inactivation in 47, XXY and 48, XXXY Klinefelter patients. Am J Med Genet 2001, 98:25–31.

24. Sanz Marcos N, Turrón Vinas A, Ibáñez Toda L: A typical presentation of Klinefelter syndrome. An Pediatr (Barc) 2012. doi:10.1016/j.anpedi.2012.10.020 [Epub ahead of print].

25. Dubois S, Illouz F, Pinson L, Bonneau D, Rohmer V, Guéchet A: Endocrine function in a 48,XXYY adult with type 2 diabetes: Case report with a review of the literature. Ann Endocrinol (Paris) 2007, 68:384–388.

26. Visootsak J, Ayari N, Howell S, Lazarus J, Tartaglia N: Timing of diagnosis of 48, XXXY and 48, XXYY: A survey of parent experiences. Am J Med Genet A 2013, 161(2):268–272.

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