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**Title:** Evidence for high breakpoint variability in 46, XX, SRY-positive testicular disorder and frequent ARSE deletion that may be associated with short stature.

**Running title:** 46, XX, SRY-positive, breakpoint variability and *ARSE*

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#### **Abstract:**

**Background:** The translocation of *SRY* onto one of the two X chromosomes results in a 46,XX testicular disorder of sex development; this is supposedly due to non-allelic homologous recombination between the protein kinase X gene (*PRKX*) and the inverted protein kinase Y pseudogene (*PRKY*). Although 46,XX *SRY*-positive men are infertile, the literature data indicate that some of these individuals are of short stature (relative to the general population). We sought to determine whether short stature was linked to additional, more complex chromosomal rearrangements.

**Methods:** Twelve laboratories gathered detailed clinical, anthropomorphic, cytogenetic and genetic data (including chromosome microarray (CMA) data) on patients with 46,XX *SRY*-positive male syndrome.

**Results:** *SRY* was present (suggesting a der(X)t(X;Y)) in 34 of the 38 cases (89.5%). When considering only the 20 patients with CMA data, we identified several chromosomal rearrangements and breakpoints – especially on the X chromosome. In the five cases for whom the X chromosome breakpoint was located in the pseudoautosomal (PAR) region, there was partial duplication of the derivative X chromosome. In contrast, in the 15 cases for whom the breakpoint was located downstream of the pseudoautosomal region, part of the derivative X chromosome had been deleted (included the arylsulfatase E (*ARSE*) gene in 11 patients). For patients with vs. without *ARSE* deletion,

the mean height was respectively  $167.7 \pm 4.5$  and  $173.1 \pm 4.0$  cm; this difference was not statistically significant ( $p=0.1005$ ).

Conclusion: Although 46,XX *SRY*-positive male syndromes were mainly due to imbalanced crossover between the X and Y chromosome during meiosis, the breakpoints differed markedly from one patient to another (especially on the X chromosome); this suggests the presence of a replication-based mechanism for recombination between non-homologous sequences. In some patients, the translocation of *SRY* to the X chromosome was associated with *ARSE* gene deletion, which might have led to short stature. With a view to explaining this disorder of sex development, whole exome sequencing could be suggested for *SRY*-negative patients.

**Keywords:** 46,XX *SRY*-positive male syndrome, replication-based mechanisms, *PRKX*, chromosomal rearrangement, short stature, arylsulfatase E gene

## Introduction

de la Chapelle et al. first described “46,XX male syndrome” in 1964 [1]. This rare disorder affects approximately one in 20,000 male infants [2,3]. As there are three phenotypes (depending on the development of male genitalia), the entity was subsequently renamed as “46,XX testicular disorder of sex development” [4]. According to the literature data, an unbalanced translocation between the X and Y chromosome during paternal meiosis [5] means that 80 to 90% of the individuals with this disorder have Y chromosomal material - particularly the sex-determining region of Y chromosome (*SRY*: chrY:2654896-2655792) gene [6]. The karyotype is then 46,X,der(X)t(X;Y)(p22.3;p11.2). Chromosome microarray (CMA) studies have shown that most breakpoints involve the protein kinase X-linked gene (*PRKX*; chrX:3522384-3631675) and the protein kinase Y-linked pseudogene (*PRKY*; chrY:7142013-7249588) [7]; these results are in line with previous studies of Y chromosome polymorphisms [8]. It has been hypothesized that the translocation results from non-allelic homologous recombination between *PRKX* and *PRKY*. In rare cases, *SRY* is located on the short arm of an acrocentric chromosome, rather than on the short arm of the X chromosome [9].

The presence of two pseudoautosomal regions (PAR1 and PAR2, located respectively at the tip of the short arm (Xp/Yp) and the tip of the long arm (Xq/Yq)) is observed in humans but not in other mammals [10,11]. Although the 2.7 Mb PAR1 region is found in most eutherian mammals and was incorporated into the sex chromosome within the last 105 million years, the 330 kb-sized PAR2

emerged after the divergence of humans and chimpanzees (i.e. approximately 6 million years ago) [12]. In PAR1, crossing-over (CO) is essential for the full disjunction of the X and Y chromosomes during male meiosis [13]. PAR1 deletion is associated with total male sterility in humans [14], and a low recombination frequency can produce aneuploid sperm [15]. Pedigree analysis has revealed that one CO per male meiosis in the PAR1 is obligatory [16]. A study of male mice showed that homologous chromosomes pair significantly later in the PAR than in the autosomes, and that this process is mediated by a different isoform of a key recombination protein (Spo11) [13]. Lastly, it has been postulated that the t(X;Y)(p22.3;p11.2) karyotype results from extension of the CO to X-specific and Y-specific regions, leading to unequal Y-to-X interchange during male meiosis. As *SRY* is the first gene downstream of PAR1 on the Y chromosome, this process leads to the translocation of *SRY* onto the X chromosome [17].

46,XX men are sterile (due to azoospermia) and can present a variety of particular physical and developmental characteristics: abnormal hair distribution, gynecomastia, low testes volume, abnormal penis size, abnormal pubic hair development, erectile function disorders, and hypergonadotropic hypogonadism. All these characteristics are associated with the X chromosome disomy and the absence of most of the Y chromosome [18]. Surprisingly, a nonnegligible proportion of “46,XX” men have short stature, relative to healthy controls and patients with Klinefelter syndrome (mean  $\pm$  standard deviation (SD) height: 170.6  $\pm$  6.0 cm, 180.7  $\pm$  6.7 cm, and 183.4  $\pm$  9.4 cm, respectively [19]) – even though the short stature homeobox (*SHOX*) gene has not been deleted (the cause of Léri-Well syndrome [20]). Similarly, with reference to both the male and female standard height curves and the target height (based on the parents' measurements), 46,XX-males are shorter and grow more slowly than expected [21]. The same study found that 47,XXY and 47,XYY individuals are taller than expected. The fact that 47,XYY boys grew slightly faster and were taller than 47,XXY boys has prompted researchers to suggest the existence of an additional, Y-specific growth gene. However, to the best of our knowledge, no such candidate genes have been identified.

We therefore hypothesized that chromosomal rearrangements leading to short stature would be more complex than expected and would affect a contiguous gene. To evaluate the putative relationship with short stature, we invited the members of a network of French cytogenetics laboratories ([www.eaclf.org](http://www.eaclf.org)) to collate their data on “46,XX” men and to characterize their patients' chromosomal rearrangements (using a CMA).

## Materials and Methods

The 37 member laboratories of the *Association des Cytogénéticiens de Langue Française* ([www.eaclf.org](http://www.eaclf.org)) who used CMAs were invited to provide all the available data (the karyotyping results, the indication for karyotyping, the age at diagnosis, height, weight, hormone levels, the presence or absence of the *SRY* gene, and the CMA results) on individuals with 46,XX male syndrome. Detailed information on each case was extracted from the corresponding medical records. We only considered “46,XX” patients for whom the karyotype indication was available and who had been tested for the presence or absence of the *SRY* gene. The CMA resolution and format varied from one laboratory to another and changed over time. All statistical analyses were performed using Statview software (version 5.0, SAS Institute Inc, Cary, NC, USA). In line with the French legislation on retrospective studies of routine clinical practice, the study protocol was approved by a hospital committee with competency for research not requiring approval by an institutional review board. All patient data were anonymized prior to analysis.

## Results

General information: Twelve laboratories accepted our invitation and supplied data on a total of 38 cases of 46,XX male syndrome diagnosed between 2001 and 2021. Karyotyping had been prompted by infertility (azoospermia) in 29 cases (76.3%), by a urogenital malformation in 6 cases (15.8%), and by another indication in 3 cases (7.9%). When considering solely adult patients with data, the mean follicle stimulating hormone and luteinizing hormone levels were respectively  $34.4 \pm 13.0$  and  $19.6 \pm 7.7$  IU/L. The *SRY* gene was detected in 34 of the 38 cases (89.5%).

CMA analysis: A CMA analysis was performed in 20 of the 34 *SRY*+ patients. We observed 14 unbalanced translocations with breakpoints on Xp22.3 and Yp11.2 (46,X,der(X)t(X;Y)(p22.3;p11.2)), one unbalanced translocation leading to a dicentric chromosome with breakpoints on Xp22.3 and Yq11.2 (46,X,dic(X;Y)(p22.3;q11.2)), two unbalanced chromosomal insertions (46,X,der(X)ins(Y;X)(p11.2;p11.2;p22.3)), and three complex chromosomal rearrangements that combined an unbalanced translocation (46,X,der(X)t(X;Y)(p22.3;p11.1)) with a deletion on Yp11.2. CMA analysis was performed in only one of the *SRY*-negative patients, and the result was normal. When taking account of the various CMA resolutions and formats, at least eight different X chromosome breakpoints were observed (Table1 and Figure 1); they ranged from the 5' region of SHOX to the 3'

region of *NLGN4*. The breakpoint was in or close to *PRKX* in 9 of the 20 patients (Figure 1 and 2). No detailed conclusions could be drawn for the Y chromosome, since the CMA was not designed to evaluate the region between *ZFY* (chrY:2,803,518-2,850,547) and *PCDH11Y* (chrY:4,924,131-5,610,264). All the breakpoints occurred downstream of *ZFY* in a 2 Mb region that lacked coding genes (Figure 3); however, none were close to *PRKY*. The breakpoint was 2,656,461 bp from the telomere in 2 cases, 2,905,059 bp from the telomere in 6 cases, 3,163,585 bp from the telomere in 9 cases, and 5,246,319 bp from the telomere in 2 cases (Table 1 and Figure 3). In the latter case, the breakpoint was located on the long arm of the Y chromosome.

**X chromosome deletion and duplication:** All 15 patients, with an X chromosome breakpoint outside the PAR1 region, had a del(X)(p22.3) on the derivate chromosome. The deletion included 1 to 7 genes. The deleted genes considered to be morbid in the OMIM database (<https://omim.org/>) were *ARSE* (coding for arylsulfatase E) in 10 cases and *NLGN4* (coding for neuroligin 4) in one case. The remaining five patients had a breakpoint in PAR1; none of these partial duplications of the X chromosome included *SHOX*.

**ARSE and height:** Given that *ARSE* inactivation in humans is associated with the X-linked recessive disease brachytelephalangi chondrodysplasia punctata (MIM #302950, *CDPX1*, associated with shortened long bones and short stature), we compared the patients' height as a function of *ARSE* deletion. Patients lacking *ARSE* (n=8) were shorter (mean height: 168.0 ± 4.7 cm) than patients bearing *ARSE* (n=7; mean height: 173.3 ± 3.8 cm) but the difference was not statistically significant (p=0.1005).

## Discussion

In line with the literature data, *SRY* was present in 80% of our cases; this observation confirmed that rearrangements between the X and Y chromosomes are the most common cause of a 46,XX testicular disorder of sex development. However, our CMA findings confirmed that although the most common rearrangement is an unbalanced translocation with breakpoints on Xp22.3 and Yp11.2, translocation breakpoints on Yq11.2 and two insertions of the Y chromosome are also possible. An additional copy number variation on Yp11.2 was identified in three cases. According to conventional karyotyping, (i) the breakpoints appeared to be similar in 17 of the 20 patients, and (ii) the X breakpoint was always on Xp22.3, regardless of the Y chromosome breakpoint. Although these findings have no implications for clinical practice, they indicate that CMA should be preferred to *SRY*-spanning FISH when seeking to better define the karyotype and chromosomal rearrangement in

male patients with a 46,XX karyotype; to avoid any incidental findings, the analysis must focus on the X and Y chromosomes only.

Unexpectedly, our CMA results demonstrated that Xp22.3 breakpoints are heterogeneous (Figure 1) – even though they were close to *PRKX* in 9 of the 20 cases. After taking account of variability in the CMA design, we identified eight different breakpoints on the X chromosome. The Y chromosome breakpoints (Figure 3) appeared to be far from *PRKY* (chrY:7142013-7249588); this finding contradicts previous reports [7,8]. Three hot spots were identified: the first close to *ZFY*, the second close to *AGPAT5P1*, and the third close to *PCDH11Y*. This result prompted us to hypothesize that a replication-based mechanism (i.e. recombination between nonhomologous sequences) is more likely than non-allelic recombination between the homologous sequences of the short arms of the X- and Y chromosomes.

In males, the CO rate in the 2.7 Mb PAR1 is approximately 17-fold greater than in the genome as a whole and over four times greater than in the next most recombinogenic genomic region of a similar size. In females, the recombination rate in PAR1 is similar to the mean genome-wide rate [22]. In contrast to PAR1 (with at least one recombination per male meiosis), it has been estimated that recombination within PAR2 occurs in just 1% of male meioses and is extremely rare in female gametogenesis [22,23]. In male PAR1, the main recombination hotspot is located within the *SHOX* gene and even constitutes one of the hottest spots in the genome [24]. However, CO activity is intense throughout PAR1, with a high frequencies close to the pseudoautosomal boundary. In females, the highest recombination rate is found near this boundary [25]. According to a study of genetic diversity in PAR1, recombination spans the pseudoautosomal boundary [23]; this would explain the occurrence of the 46,XX testicular disorder of sex development and then the breakpoint variability observed in the present series.

Twelve of the 20 patients had an X chromosome deletion, and eleven of the 15 deletions including *ARSE* (Figures 1 and 2). As mentioned above, *ARSE* inactivation can lead to CDPX1. Given that *ARSE* is the only OMIM-listed gene recurrently deleted from the X chromosome, we hypothesize that *ARSE* deletion is associated with the short stature in “46,XX” males. Females lacking *ARSE* have not been described clinically but some women are of short stature [26]. For a few X-linked diseases, female carriers are expected to have skewed X-chromosome inactivation (sXCI) and clinical findings [27][28]. No data are available for female carriers of CDPX1, since the prognosis for men with CDPX1 is quite good; however, some affected men have significant medical problems, including respiratory compromise, cervical spine stenosis and instability, mixed conductive and sensorineural hearing loss, and intellectual disability [26]. We therefore hypothesize that stature depends on sXCI in female carriers. Some of the patients in our series probably had sXCI, since the rearranged X chromosome



had been predominantly activated and the male phenotype was still normal. The mean height in our study population ( $170.4 \pm 6.1$  cm) is quite similar to a value of  $170.6 \pm 6.0$  cm reported previously for 46,XX males [19]. In our series, the patients with *ARSE* deletion were shorter than those still bearing *ARSE* (mean  $\pm$  SD height:  $168.0 \pm 4.7$  vs.  $173.3 \pm 3.8$  cm, respectively) but the difference was not statistically significant ( $p=0.1005$ ). However, no firm conclusions can be drawn for the 15 patients (including 8 without the *ARSE* deletion), even though they were shorter (on average) than the French male population in 2019 (mean height: 177.8) [29]. Regardless of the real impact of the *ARSE* deletion, 46,XX testicular disorder of sex development should be considered as a particular type of contiguous genetic syndrome that combines sex reversal and short stature. The sex reversal is due to the *SRY* translocation on the X chromosome, and the short stature is probably due to the *ARSE* deletion.

### Conclusion

Chromosomal rearrangements in 46,XX,SRY+ male patients appear to be more complex than originally thought; in particular, they are highly variable with regard to the X chromosome breakpoint. Moreover, many of these chromosomal rearrangements are combined with *ARSE* deletion, which might explain the short stature observed in some patients and correlate the phenotype to the genotype. Further characterization of 46,XX males is required to confirm this hypothesis and thus to establish whether the 46,XX testicular disorder of sex development is a contiguous gene syndrome in some cases. A major limitation of efforts to correlate *ARSE* deletion with the patient's height is the absence of data from whole genome sequencing, whole exome sequencing, and/or RNA sequencing. If pathologic variants cannot be ruled out, no clear conclusions can be drawn.

**Data Availability Statement :** Yes

### Author contribution statement

Patient inclusion: SJ, PK, GS, GN, PC, MPB, BH, JMD, NG, PC, FA, ED, PV, IP, CC, MABR, CSB, FV

CMA analysis: LJ, GV, SJ, PK, GS, BH, JMD, RH, CSB, FV

Study design: FV

Writing: CC and FV

Proofreading: LJ, GV, SJ, PK, GS, IP, CC, CSB

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### Figure legends:

Figure 1: X chromosome rearrangements with the location of *ARSE* in light blue. A to C: the blue line corresponds to partial duplication. D to H: the red line corresponds to partial deletion. Specific breakpoint locations are given in Table 1.



Figure 2: CMA results, using male (A/B, E/F and G/H) or female (C/D) DNA references.

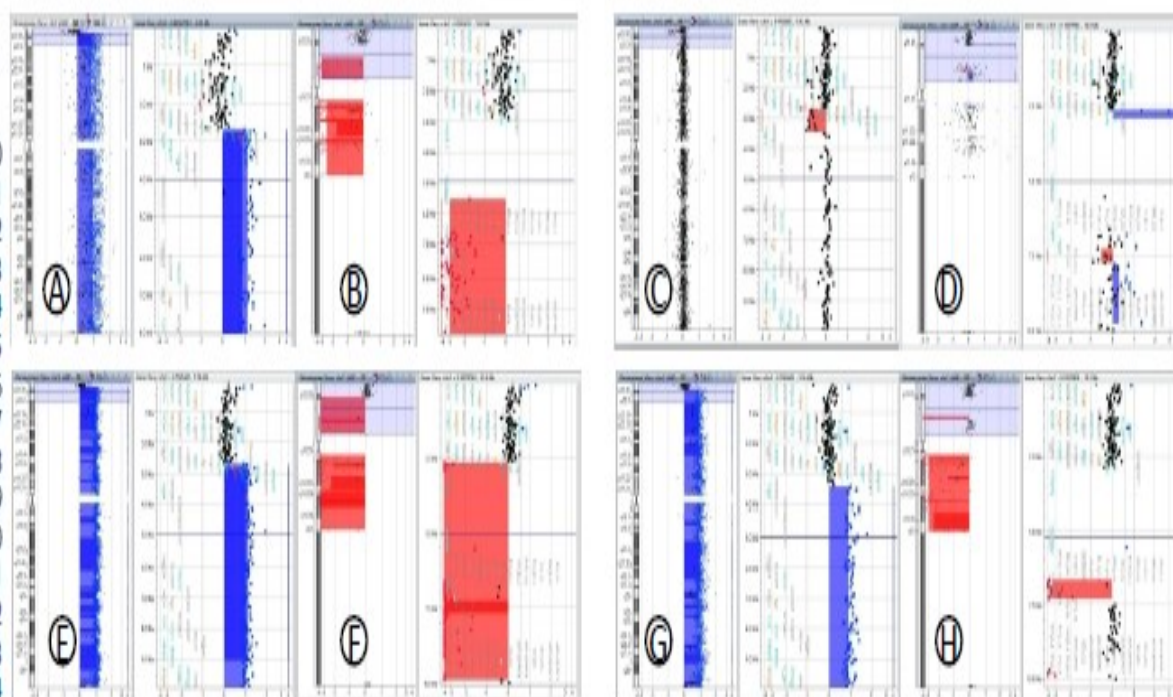
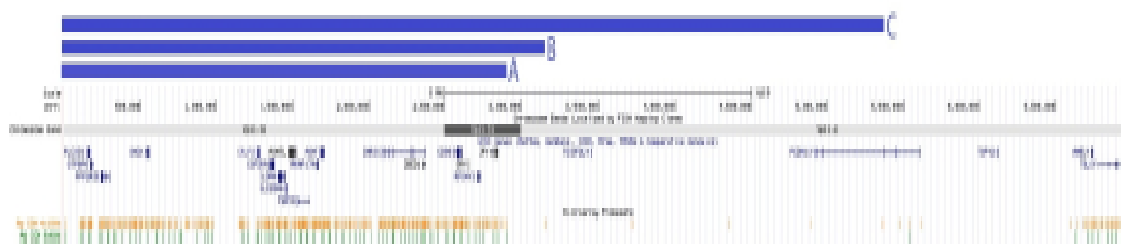


Figure 3: Y chromosome rearrangements. A to C; the blue line corresponds to partial duplication. Specific breakpoint locations are given in Table 1.



**Table legend:**

Table 1: Detailed chromosome microarray data for the 20 “46,XX,SRY positive” patients.

Patient number	Karyotype interpretation	Agilent array format	Chromosome X abnormality (1,838,587)	X chromosome breakpoint	X chromosome rearrangement type	Chromosome Y short arm abnormality	Y chromosome short arm breakpoint	Y chromosome rearrangement type
1	46,X,der(X)t(X;Y)(p22,3;p11,2)	18 Ok	2,689,408 x3	1,838,587	B	(1-3,163,585)x1	3,163,585	B
2	46,X,der(X)t(X;Y)(p22,3;p11,1),del(Y)(p11,1)	18 Ok	3,590,348 x1	3,590,348	F	(1-3,163,585)x1,(7,184,216-9,458,094)x1	3,163,585	B
3	46,X,der(X)t(X;Y)(p22,3;p11,2)	18 Ok	2,689,408 x3	2,120,345	C	(1-3,163,585)x1	3,163,585	B
4	46,X,der(X)t(X;Y)(p22,3;p11,2)	18 Ok	3,548,012 x1	3,548,012	F	(1-3,163,585)x1	3,163,585	B
5	46,X,der(X)t(X;Y)(p22,3;p11,2)	18 Ok	3,469,500 x1	3,469,500	E	(1-3,163,585)x1	3,163,585	B
6	46,X,der(X)t(X;Y)(p22,3;p11,2)	18 Ok	2,689,408 x3	1,789,694	B	(1-2,905,059)x1	2,905,059	A
7	46,X,der(X)t(X;Y)(p22,3;p11,2)	18 Ok	(2,701,273)	6,182,748	H	(1-2,905,059)x1	2,905,059	A

				6,182,748)					
				x1					
				(2,701,273					
				-					
8	46,X,dic(X;Y)(p22,3;q11,2)	18 Ok	3,524,184)	3,524					
			x1	,184	F	(1-9,650,194)x1	/	/	
			(2,701,273						
9	46,X,der(X)t(X;Y)(p22,3;p11,2)	18 Ok	-	2,716,319)				3,163	
			x1	,319	D	(1-3,163,585)x1	,585		B
			(2,701,273						
10	46,X,der(X)t(X;Y)(p22,3;p11,2)	18 Ok	-	3,351,301)				5,246	
			x1	,301	E	(1-5,246,319)x1	,319		C
			(697,227-						
11	46,XX,ins(Y;X)(p11,2p11,2;p22,3)	18 Ok	2,689,408)	697.2				3,163	
			x3	27	A	(1-3,163,585)x1	,585		B
			(2,709,027						
12	46,X,der(X)t(X;Y)(p22,3;p11,2)	60 k	-	3,469,559)				2,905	
			x1	,500	E	(1-2,905,059)x1	,059		A
13	46,X,der(X)t(X;Y)(p22,3;p11,2)	60 k	/	/	/	(1-2,905,059)x1	,059		A
			(2,709,027						
14	46,X,der(X)t(X;Y)(p22,3;p11,1),del(Y)(p11,1)	60 k	-	3,376,702)				2,905	
			x1	,702	E	(1-2,905,059)x1,(7,5	12,591-	2,905	
			(2,701,273			9,394,173)x1	,059		A
15	46,X,der(X)t(X;Y)(p22,3;p11,2)	18 Ok	-	5,395,569)				3,163	
			x1	,569	G	(1-3,163,585)x1	,585		B
			(1,755,742						
16	46,XX,ins(Y;X)(p11,2p11,2;p22,3)	18 Ok	2,689,408)	1,755				2,905	
			x3	,742	B	(1-2,905,059)x1	,059		A
			(2,701,273						
17	46,X,der(X)t(X;Y)(p22,3;p11,1),del(Y)(p11,1)	18 Ok	-	3,351,301)				5,246	
			x1	,301	E	(1-5,246,319)x1,(7,5	30,569-	,319	
			(2,701,273			9,458,094)x1			C
18	46,X,der(X)t(X;Y)(p22,3;p11,2)	18 Ok	-	2,716,319)				2,656	
			x1	,319	D	(1-2,905,059)x1	,461		A
			(2,701,273						
19	46,X,der(X)t(X;Y)(p22,3;p11,2)	18 Ok	-	2,716,319)				2,656	
			x1	,319	D	(1-2,905,059)x1	,461		A
			(2,701,273						
20	46,X,der(X)t(X;Y)(p22,3;p11,2)	18 Ok	3,524,184)	3,524				3,163	
			x1	,184	F	(1-3,163,585)x1	,585		B

X chromosome abnormality: only partial duplications of the pseudoautosomal region are shown (cases 1, 3, 6, 11 and 16), or deletions after the pseudoautosomal region.

The types of X chromosome breakpoints are linked to Figure 1

Y chromosome abnormality: The complete sequence of the Y chromosome included in the derivative chromosome is described, including the pseudoautosomal region.

The types of Y chromosome breakpoints are linked to Figure 3