CLINICAL AND CYTOGENETIC STUDY OF TURNER SYNDROME AND ITS VARIANTS

Shailaja CM 1, Shobha *2, Vijayakumar BJ 3, Pravinkumar NK 4.

1 Professor, Department of Anatomy, SSIMS & RC, Davangere, Karnataka, India.
2 Associate Professor, Department of Anatomy, JJMMC, Davangere, Karnataka, India.
3 Professor & Head, Department of Forensic Medicine & Toxicology, SSIMS & RC, Davangere, Karnataka, India.
4 Associate Professor, Department of Forensic Medicine & Toxicology, SSIMS & RC, Davangere, Karnataka, India.

ABSTRACT

62 cases of Turner syndrome (T.S) were clinically diagnosed and sent to cytogenetic laboratory for confirmation of the diagnosis through karyotyping from 2012 to 2018. Out of 62 cases, 43 turned out to be T.S & its variants. Most commonly observed karyotype was 45,X (62.79%), followed by 45,X/46,XX (23.25%), 45,X/46,Xi (6.97%) mosaicism, 4.65% of 46,Xi & 2.32% of a very rare type of variant of T.S i.e 45,X/46,XY.

Patients with 45,X karyotype had typical features of T.S such as short stature (< 5feet or 150cms), delayed appearance of secondary sexual characters & dysmorphic facies, the main complaint of these patients was primary amenorrhoea. Patients were younger at diagnosis & had a significant shorter mean adult height than those with 45,X/46,XX mosaicism. Those with mosaicism had mild dysmorphic features & presented with primary or secondary amenorrhoea. The rarest type of T.S (45,X/45,XY) was presented in a new born with ambiguous genitalia (suspected for CAH).

Short stature with sexual infantilism & primary or secondary amenorrhoea in a young female should suggest the possibility of Turner syndrome, which should be confirmed by chromosome analysis.

KEY WORDS: Turner Syndrome and Variants, Cytogenetics.

INTRODUCTION

Turner syndrome is the consequence of complete or partial absence of one X chromosome in a phenotypic female usually characterized by short stature, gonadal dysgenesis & a variety of other features.

In 1930, Otto Ullrich had reported on an 8 yrs old girl with short stature, lymphedema web neck, cubitus valgus & dysmorphic features, which was the 1st case of T.S [1] But it was 1st described as a distinct entity by Turner in 1938, in 7 females with short stature & sexual infantilism, webbed neck & cubitus valgus & dysmorphic features [2].

Patients with 45,X/46,XY mosaicism exhibit a wide phenotypic spectrum ranging from normal male (90%), which go unnoticed till puberty or...
present with hypospadiasis to male or female pseudo hermaphroditism. 10% females present with incomplete male development & appear as partly masculinised female genitalia. The term Mixed Gonadal Dysgenesis (MGD)/ Disorder of Sex Development (DSD) is sometimes used for this group. Telvi et al in 1999 [3] reviewed post-natal ascertained 45,X/ 46,XY cases in which some males who appeared normal at birth can develop late onset “Turner syndrome like” abnormalities. New classification proposed by Chicago consensus (Hughes IA et al 2006 & Lee et al 2006) [20] the term MGD has been employed in cases of testicular dysgenesis with 45,X/ 46,XY karyotype & partial gonadal dysgenesis (PGD) in those with 46,XY constitution, regardless of histological picture.

Patients with Y mosaicism have high risk for developing gonadoblastoma & dysgerminoma. Here the main aim of our study is to co-relate the clinical profile with that of karyotypic abnormalities in a T.S & its variants.

MATERIALS AND METHODS

62 cases of clinically diagnosed T.S were sent to the cytogenetic laboratory (SSIMS & RC Davangere) for karyotyping from 2012 to 2018. All the cases were analysed & evaluated prospectively.

Their presenting signs & symptoms, clinical features & the age of presentation were included. Hormonal profile (Serum FSH, LH & Serum Electrolytes in special cases) & pelvic ultrasound findings were also included in the study. A detailed evaluation of CVS was considered in suspected cases.

Peripheral blood lymphocyte culture was carried out by standard technique of Moorhead et al [3,5]. Trypsin & Giemsa or G banding of metaphase chromosomes was carried out. 25 metaphases of each patient were analysed, if any one metaphase showed a different arrangement of chromosomes then another 25 metaphases were analysed. If the different cell line was noted again then another 25 metaphases were analysed with a total of 75 metaphases. This was done to rule out mosaicism. Serum FSH, LH were analysed in standard laboratory with ELISA method. In one of our case with ambiguous genitalia (suspected for congenital adrenal hyperplasia), serum electrolytes were also analysed.

The mean age (>16 years) & mean adult height (< 5 feet/150 cms) at the time of diagnosis were compared with 45,X & 45,X/ 46,XX mosaic group. These features were not compared with third group which showed heterogenous mix of chromosomal abnormalities associated with T.S.

RESULTS

Table 1: Cytogenetic abnormalities observed in T.S are shown below.

<table>
<thead>
<tr>
<th>Normal Number</th>
</tr>
</thead>
<tbody>
<tr>
<td>46,XX</td>
</tr>
<tr>
<td>CLASSIC</td>
</tr>
<tr>
<td>45,X</td>
</tr>
<tr>
<td>MOSAICISM</td>
</tr>
<tr>
<td>45,X/ 46,XX</td>
</tr>
<tr>
<td>OTHER/ THIRD GROUP</td>
</tr>
<tr>
<td>45,X/ 46Xi(Xq)</td>
</tr>
<tr>
<td>46,Xi(Xq)</td>
</tr>
<tr>
<td>45,X/ 46,XY</td>
</tr>
<tr>
<td>TOTAL</td>
</tr>
</tbody>
</table>

Monosomy X (45,X): Out of 27 cases with 45,X karyotype ,18 patients were d”16yrs, including a new-born (2days old) . Other 9 patients were above the age of 16.The mean age at the time of diagnosis was 15yrs. The mean adult height was 123.6 cms (Table 2). The presenting complaint of 18 patients was short stature with some physical findings as shown in Table 2.The new born presented with ambiguous genitalia. Other 9 patients presented with primary amenorrhoea as their main complaint.

45,X/ 46,XX Mosaicism: Out of 10 patients in this group ,7 were of >16 yrs of age while other 3 were d” than 16yrs of age. The mean age at diagnosis was 19 yrs & the mean adult height was 152.2 cms.

In 7 patients main presenting complaint was primary amenorrhoea, in other 3 patients – one presented with secondary amenorrhoea, other 2 presented with short stature & delayed sex characters. All other Turner stigmata were infrequently seen (Table 2).

Other X chromosome abnormalities associated with Turner syndrome: Apart from 45,X & 45,X/ 46XX mosaicism other cytogenetic abnormalities seen are shown in Table 1.
Out of 6 patients, 3 patients showed 45,X/45Xi (Xq), 2 patients showed 46,Xi (Xq) & 1 patient (new born) showed 45,X/46,XY.

5 patients of this group were > 16 yrs of age. Mean age at diagnosis was 19 yrs of age & mean adult age was 156cms. Out of 5 patients 3 presented with primary amenorrhoea with delayed sex characters & 2 patients with secondary amenorrhoea. Other physical features are as shown in Table 2.

The new born with mosaicism for Y chromosome suspected for CAH was included in our study as it had presented with ambiguous genitalia (external genitalia showed a large phallus of 1.3cms, with no separate urethral opening, urogenital sinus with wide opening, scrotal sac with rugocity seen, but no contents, no hernia in inguinoperineal region seen) & had a characteristic Turner phenotype.

**Hormonal profile:** Estimation of serum FSH & LH could be elicited in only 21 patients out of 34 who were above the age of 15 yrs. This includes 9 patients in 45,X Monosomy, 7 patients in 45,X/46XX mosaicism & 5 patients in other group of karyotype. The mean FSH & LH levels were 55.6 ± 45.4 (range -1.8-172) & 18.3 ± 15.1 (range 0.8-51.5) mlU/ml, respectively. These levels were not compared between these three groups, as the number of patients in each type was very small.

As the new born was suspected for Congenital Adrenal Hyperplasia (CAH), serum electrolytes were analysed (serum sodium 130.5mmol/l, serum potassium 5.5mmol/l & serum chloride 100.1mmol/l), all were within normal limits, & were estimated for 3 consecutive days.

Thyroid function tests were carried out in 12 patients, who came with main complaint as short stature. Out of 12 patients, 3 of them showed elevated TSH & low thyroxin levels suggesting early hypothyroidism. They were referred to endocrinologist for further evaluation.

**Pelvic ultrasound:** Pelvic ultrasound findings were available for 20 out of 34 patients who were more than 15 yrs of age. These comprised of 9 patients with 45,X, 6 patients with 45,X/46XX & 5 patients from other group of karyotype. 50% of these patients showed hypoplastic uterus with bilateral streak ovaries. 35% of them showed unilateral or bilateral absence of ovaries & 15% of them showed normal uterus & ovaries. While the new born with ambiguous genitalia showed presence of uterus (2.7cms x 9mm x 1.5cms) with normal endometrium. No suggestions of testis like structure seen in inguinoperineal region. Kidneys & adrenal glands were normal in size & echo-texture.

**Table 2:** Clinical co-relation with karyotype in T.S & its variants are shown in table 2. (percentage).

<table>
<thead>
<tr>
<th>Features</th>
<th>45X</th>
<th>45,X/46XX</th>
<th>Others</th>
</tr>
</thead>
<tbody>
<tr>
<td>Short stature</td>
<td>90</td>
<td>75</td>
<td>59</td>
</tr>
<tr>
<td>Dysmorphic face</td>
<td>50</td>
<td>10</td>
<td>10</td>
</tr>
<tr>
<td>Short neck</td>
<td>10</td>
<td>5</td>
<td>5</td>
</tr>
<tr>
<td>Low posterior hair line</td>
<td>5</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>Cubitus valgus</td>
<td>25</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>Shield chest</td>
<td>35</td>
<td>5</td>
<td>4</td>
</tr>
<tr>
<td>Delayed sex characters</td>
<td>100</td>
<td>75</td>
<td>56</td>
</tr>
<tr>
<td>Primary amenorrhoea</td>
<td>100</td>
<td>65</td>
<td>50</td>
</tr>
<tr>
<td>Secondary amenorrhoea</td>
<td>0</td>
<td>30</td>
<td>20</td>
</tr>
<tr>
<td>Congenital heart disease</td>
<td>0</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>I.Q/ mental ability</td>
<td>5</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>Mean age at diagnosis (years)</td>
<td>15</td>
<td>19</td>
<td>19</td>
</tr>
<tr>
<td>Mean adult height (cms) at diagnosis</td>
<td>123.6</td>
<td>152</td>
<td>156</td>
</tr>
</tbody>
</table>

**DISCUSSION**

The approximate incidence of Turner syndrome is estimated to be 1 in 2500 live female births [8]. 45,X cell line occurs as a result of non-disjunction at either stage of meiosis 1 or 2 during spermatogenesis or oogenesis or from postzygotic error.

It is suggested that a gene dosage inequality due to absence of part or all of the X chromosome is responsible for the phenotype [9-11].

Prior to 12 weeks of in-utero development, the ovaries in 45,X female appear normal histologically, but thereafter, there is a decrease in the number of follicle cells per oocyte. In the absence of second X chromosome, the oocytes degenerate more rapidly than normal, so that at the time of birth there are few, if any left, the ovarian tissue resembles fibrotic streaks [11]. The streak gonads appear identical regardless of karyotype [6].

Short stature & primary amenorrhoea were the commonest presenting complaints of our patients, regardless of karyotype. The final adult height was significantly less in patients with...
45,X karyotype when compared with 45,X/46XX mosaicism (p< 0.05). Other common features like dysmorphic facies, cubitus valgus, shield chest (table2) are seen in 45,X group & infrequently in 45,X/46XX & less frequently in third group.

Few of the patients with 45,X/46XX mosaicism had secondary amenorrhoea (Table -2). These patients had low levels of mosaicism i.e more of 46,XX than 45X. Between 10-20% of the patients with T.S have spontaneous onset of menstruation [11]. This is seen less often in 45X group. Pregnancy was reported in 2 of our patients, one with 45,X/46,XX & other patient with Isochromosome 46,Xi(Xq). Pregnancy is rare in patients with 45,X group. Approximately more than half of the patients of T.S are likely to have miscarriages, foetal malformations or chromosomal abnormalities [7].

Mental ability was normal in our patients, only 2 patients with 45,X had IQ less than 60%. A much higher incidence of mental disability has been reported in patients with a ring chromosome [12].

Only 2 of our patients had congenital cardiovascular anomaly. Both females belonged to 45,X group & had atrial stenosis. The incidence of congenital heart disease in Turner syndrome is reported to be highest in 45,X group (10-15%) & lowest in other group (0-8%) comprising long or short arm deletions of X chromosome [11,10,4,6].

2-5% of Turner syndrome patients show mosaicism for Y chromosome. We had a new born with 45,X/46,XY karyotype & had a characteristic Turner phenotype. Traditionally these patients are said have mixed gonadal dysgenesis.

Dysgenetic testis may be bilateral or associated with contra lateral streak in subjects with 45,X/46XX karyotype. Histologically, they may show fibrous tissue & few tubular structures with reduced germ cells. As a consequence ,the external genitalia ranges from male to female, including cases of striking genital ambiguity & persistence of Mullerian structure [19].

Patients with Y chromosome are at high risk for developing gonadoblastoma [14]. Recently, molecular methods have been used to demonstrate the SRY gene in these patients, as a marker for Y chromosome [15].

CONCLUSION

Turner syndrome is a chromosomal disorder which is frequently misdiagnosed or missed completely. The severity of the syndrome decreases as it becomes mosaic. Hence any female of short stature with primary or secondary amenorrhoea, even with the absence of other phenotypic features of T.S should be suspected as T.S but confirmed by chromosomal analysis (karyotyping).

Abnormal maternal serum levels & anomalies of kidney & uterus during 2nd trimester should
warn for karyotyping. Early recognition of T.S & timely investigations will help in improving the quality of life, by improving the adult height in those responding to growth hormone therapy & initiating sex hormone replacement. Genitoplasty & gonadectomy should be preferred in those with y chromosomal material, as they are at risk for developing gonadoblastoma or dysgerminoma later in life. Such patients are asked for follow up, to monitor their growth & puberty.

Conflicts of Interests: None

REFERENCES