

**Introduction**

Turner syndrome (TS) is a genetic disorder that was first described by Turner (1) in 1938, and is the result of the complete or partial absence of the X chromosome. TS is one of the most common chromosomal abnormalities among female live births with estimated frequency of 1/2500 (2). Clinical findings include short stature, primary amenorrhea, infertility and characteristic stigmata. The phenotypic features are low nuchal hairline, low-set ears, small mandible, cubitus valgus, nail hypoplasia, high palate, swelling of the hands and feet in the neonatal period, short 4th metacarpal bone, discrete nipples and wide thoracic cage. Cardiac and renal anomalies, hypothyroidism, hearing and vision disorders, gastrointestinal and dermatological problems and neoplasms may also be seen in TS (3). Although some of the patients may be diagnosed at birth due to the presence of dysomorphic findings, diagnosis is delayed until childhood, adolescence or later. Intelligence is generally not affected. However, learning difficulties that affect nonverbal, perceptual, motor and visuo-spatial skills can be seen in 70% of the patients (4). In a recent study the authors compared TS patients with healthy individuals and suggested that verbal and nonverbal cognitive impairment may be due to anomalies in gray matter development (5).
Approximately 50% of TS patients have 45,X karyotype and the most common structural X chromosome abnormality is isochromosome Xq. The retained X chromosome is maternally derived in two thirds of the patients. The patients with 46,XY cell line mosaicism or structural rearrangement of the Y chromosome mostly have masculinized external genitalia and are at increased risk of developing gonadoblastoma and other gonadal tumors, whereas patients with mosaic 46,XX karyotype or isochromosome Xq have a milder phenotype (6).

TS with 45,X karyotype has been observed in 1-2% of human conceptions, 10% of first trimester pregnancy losses and 1% of stillbirths. More than 99% of 45,X fetuses end with abortion, typically by the 28th week of gestation, which suggests that living 45,X individuals must have mosaicism for another cell line (7,8). The etiology includes pre and post-conceptional errors, as well as meiotic non-disjunction and anaphase lag.

The pseudoautosomal regions (PAR) of X and Y chromosomes are both necessary for normal development. It has been reported that haplo-insufficiency in the CSF2RA gene located in the PAR1 region, which plays a role in normal placental development, is responsible for high mortality in 45,X karyotype embryos (9,10). Recently, it has been suggested that all living patients in whom TS with 45,X karyotype has been diagnosed, are cryptic mosaics. Loss of X chromosome in these embryos frequently result in mitotic error, but it has been reported that this loss may occur at various stages of postfertilization (11).

The diagnosis is made by karyotype analysis of peripheral blood which shows the numerical and/or structural abnormalities of the X chromosome. However, mosaicism can be better defined in approximately 30% of non-mosaic patients by increasing the number of cells examined in the karyotype analysis, evaluating additional tissue samples or using fluorescence in situ hybridization method (12).

In this review, we aimed to evaluate the phenotypical characteristics, clinical variability and the responsible genes of TS and its variants, and provide a general approach to patients with suspected TS.

Turner Syndrome with Structural Abnormalities of the X Chromosome

Isochromosome X [46,X,i(Xi)]

Isochromosome is a structural chromosomal aberration consisting of two short arms or two long arms, which are derived by centromere division. The most common structural abnormality of X chromosome is 46,X,i(Xq). The frequency of isochromosome was reported to be 15-18% in TS cases with or without mosaicism (13). Short stature, which is one of the most frequently observed clinical features, can be explained by SHOX haplo-insufficiency (14). Therefore, while short stature is more frequently present in patients with 46,X,i(Xq) karyotype, gonadal dysfunction is more likely to be seen in patients with 46,X,i(Xp) karyotype, which is a very rare entity.

The incidence of autoimmune thyroid disease in TS increases with advanced age. Recently, a twofold increase in the prevalence of autoimmune thyroid disease has been observed from the first to the third decade of life in patients with the isochromosome karyotype (15). In addition, patients with 45,X or 46,X,i(Xq) karyotype carry a higher risk of developing hearing loss when compared to patients with mosaic karyotypes. A linear relationship was also found between age and hearing loss (16).

Ring Chromosome [46,X,r(Xi)]

Ring chromosomes usually result from two terminal breaks in both chromosome arms, followed by the fusion of the broken ends. The asentric part often disappears and partial monosomy occurs. Ring chromosomes cause complex mitotic events. The phenotype is highly variable according to the size of ring chromosome and the deletions of short and long arms. A ring X chromosome is found in approximately 6% of the patients with TS, generally with mosaicism for a 45,X cell line (4).

Females with 46,X,r(X) karyotype may have typical TS findings such as short stature, peripheral edema, characteristic facial features, low neck hairline, ovarian dysgenesis, and endocrine disorders. On the other hand, mental retardation, learning disability, autism spectrum disorders, and structural brain abnormalities are more frequently observed in TS with ring chromosome than TS with 45,X karyotype. Some patients may have more severe phenotypical features and this is thought to be related to whether the X inactivation center is functional or not (17).

In normal females one of the two X chromosomes becomes inactive in the early stages of life and the expression of the X-linked genes is equalized for both sexes. The inactivation of an X chromosome is random, which is maintained in a clonal manner throughout subsequent cell divisions. As a result, females become mosaic in terms of gene expression associated with the X chromosome (18).

X inactivation center, which plays an important role in the expression of the X-inactive specific transcript (XIST) gene, is localized on the X chromosome at position q13. Also, XIST gene appears to be a key master regulatory locus for X inactivation. In patients with structural abnormalities of the X chromosome, the abnormal X chromosome (such as deletions, duplications, isochromosomes) is always the one inactivated. Smaller ring chromosomes may lack the XIST locus, rendering them functionally disomic for the genes present on the ring. Females with mosaic ring X chromosome without XIST expression can also have extremity anomalies, abnormal pigmentation and facial features of Kabuki make-up syndrome in addition to mental retardation (18,19). Hyperinsulinemic hypoglycemia is an atypical feature which can be seen especially in patients with mosaic karyotype (20).

Deletion (Xp or Xq)

The frequency of Xp deletion in patients with TS is approximately 2% (4). Short stature, gonadal dysgenesis
and characteristic TS stigmata are especially observed in patients who show deletion of the entire short arm. Furthermore, the phenotype is variable in partial deletions. The region Xp22.33-Xp22.12 contains the SHOX gene, which is located in the terminal region. The gene escapes from X inactivation, and its function is dosage dependent. Therefore, haplo-insufficiency of SHOX gene causes growth retardation (21). The SHOX gene is expressed in the pharyngeal arch, limbs, osteogenic cells, bone marrow, and fibroblasts. As well as short stature, skeletal abnormalities including short metacarpals, high palate, cubitus valgus, Madelung deformity, and mesomelic dysplasia may also be seen in the haplo-insufficiency of the gene. (22). Mutations involving the SHOX gene or its regulatory regions can be detected in approximately 17% of the patients with idiopathic short stature, and in 50-90% of the patients with Leri-Weill syndrome (23).

The genes located on Xp and Yp are necessary for normal development of the cardiovascular system. Congenital heart diseases have an important role in prenatal and postnatal mortality in patients with TS. The most common cardiovascular defects are bicuspid aortic valve (30%) and aortic coarctation (12%). Additionally, hypoplastic left heart syndrome, usually lethal, is seen in about 10% of the patients (24,25). In particular, left ventricular outflow tract defects are associated with the terminal region of Xp.

The cytogenetic studies have shown that the region between Xq13 and Xq28 is important for normal ovarian function. The Xq13-q21 region is defined as the critical region (CR). The proximal deletions of this region are usually compatible with normal menstruation and fertility. The terminal and interstitial deletions of the CR2 at Xq23-q28 are mostly responsible for premature ovarian failure (26). The FMR1 gene, which is important in the ovarian function, is localized in the Xq27.3 locus, and expansions in the exon 1 triplet repeat of the gene are associated with an increased risk of early menopause (27).

**Turner Syndrome with Mosaicism**

Although classic TS karyotype is 45,X, 30-40% of the remaining have a mosaic pattern with a second cell line (45,X/46,XX, 45,X/47,XXX, 45,X/46,XY, 45,X/47,XYY and 46,X,del(Xq)) (4,28).

**45,X/46,XX**

The most common form of mosaicism is 45,X/46,XX (15%) which includes both the cell line with the normal karyotype and the pathological cytogenetic structure (29). These patients may have a normal phenotype rather than typical TS features. However, spontaneous menstruation occurs in about 3% of the 45,X females, and up to 20% of mosaic females. While the mean adult length is higher in mosaic females, the probability of the presence of a somatic anomaly is lower.

Psychiatric disorders can also be seen. Especially, the majority of TS patients with schizophrenia had a mosaic 45,X/46,XX karyotype. It has been suggested that the potential of gene dose-effect might be associated with the abnormal expression of an X chromosome gene product which has a susceptibility for schizophrenia in TS (30). Recently, it has been suggested that the HOPA gene which is located on Xq13, might be involved in the development of schizophrenia as well as being associated with mental retardation and thyroid dysfunction (31).

**45,X/46,XY**

TS with 46,XY cell line can be seen in 5-10% of the patients (32). Its clinical features are quite variable. The patients may have typical TS phenotype, normal male appearance, moderate masculinization, male pseudohermaphroditism, and mixed gonadal dysgenesis which can be transformed into a malignant form (33). However, the presence of Y cell line cannot be predicted from the phenotype. Patients with a normal female phenotype without evidence of masculinization may also have a 46,XY cell line. Therefore, the patients should be closely followed-up due to the risk of any germ cell tumor, especially gonadoblastoma. In current literature, the frequency of developing germ cell tumor is reported to be about 15% in patients with 45,X/46,XY karyotype (32).

**45,X/47,XXX**

Triple X syndrome occurs at a frequency of 1/1000 in live-born females. Clinical features include long stature with large hands/feet and microcephalic appearance, hypotonia, seizures, genitourinary abnormalities, and premature ovarian failure. Onset of puberty, sexual development, and fertility are usually normal. Speech and motor retardation, learning disability, attention deficit and behavioral disorders may be seen also in patients with 47,XXX karyotype (34).

The 45,X/47,XXX karyotype, which is extremely rare, constitutes 3-4% of the patients with TS and the etiology can be attributed to post-zygotic non-dysjunction that occurs in the normal disomic cell lines. The patients usually have a milder phenotype and the characteristic stigmata of TS may not be seen. Ovarian failure may be observed however, and spontaneous menarche and fertility are more common in patients with 45,X/47,XXX karyotype than those with monosomy (35). Previous studies reported the frequency of spontaneous menarche and fertility as 84% and 69% respectively in patients with 45,X/47,XXX karyotype (36). Moreover, there was no significant difference between the patients with 45,X karyotype and those with 45,X/47,XXX and 45,X/46,XX/47,XXX karyotype with regard to the development of mental retardation. However, cell counts that provide a ratio of 45,X to 47,XXX cells should not be considered to have predictive value, because they vary in different tissues (36,37).

In conclusion, TS is a complex, reproductive and developmental disorder. Because the clinical features are
quite variable, the phenotype-genotype correlation may not always be achieved. For this reason, karyotype analysis should be done in patients with a clinical suspicion of TS, even in the absence of the phenotypic stigmata, and the probability of a mosaic karyotype should be kept in mind. Early diagnosis will improve the life quality of these patients and could prevent future problems that may arise during adulthood.

Ethics

Peer-review: Internally peer-reviewed.

Authorship Contributions


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