Introduction

Infertility, the inability to conceive by natural means, is a worldwide problem affecting 8-12% couple during their reproductive lives with high prevalence (10-15%) in India. It is estimated that while female factor accounts for 40-50% of infertility among couples, infertility attributable to male factors is on the rise and constitutes 30-40% of infertility [1,2]. Infertility can be hormonal, related to age, exercise, obesity or infectious disease; it can be immunological, psychological, result from surgery or blockage, or be associated with defined abnormalities in the gametes. Many factors are implicated in the etiology of infertility, be it male associated or attributed to the female partner. These factors may be hormonal, infectious, immunological, surgical or psychological. Most of these factors have genetic basis involving several genes and gene products. In the future, pursuing the most promising genetic variants, mutations, or polymorphisms may provide clinically relevant therapeutics for infertile individuals. As more genes are discovered and the etiology of infertility disorders becomes well understood, the management and treatment of infertility will improve as well. Herein, the review presents the known genetic causes and their associations for both male and female infertility.

Male Infertility

Structural Chromosomal Abnormalities (SCAs)

It includes deletions, duplications, translocations (balanced, imbalanced and Robertsonian) and inversions. There are two alternative models that explain the aberration effect. First, it blocks spermatogenesis via abnormal chromosome synapsis in crossover and meiosis arrest. Second, the aberration disrupts a dosage-sensitive gene, resulting in spermatogenesis arrest and infertility [3].

Translocations

Chromosomal translocations can be of two types, i.e., Robertsonian translocations involve acrocentric chromosomes (13, 14, 15, 21 and 22) and reciprocal translocations involve mutual exchange of chromosomal segments between autosomal and sex chromosomes. Chromosomal translocations may cause reductions in testicular volume and testosterone level, which may impact spermatogenesis, resulting in azoospermia or oligozoospermia and thereby, male infertility [4,5].

Y Chromosome Deletions

It is estimated that 19% of males diagnosed with idiopathic infertility have Yq (long arm) micro-deletions and between 50-70% of the non-recombining region of human Y chromosome is composed of a variety of highly repeated DNA elements, the majority of which appear to be unique to the human Y chromosome. Deletions of the Y chromosome are likely to be consequence of these repeated elements causing intra-chromosomal recombination. There is a possibility that Y chromosome micro-deletions may also contribute to spermatogenic failure. The type and severity of structural anomalies depends on the location and size of the anomaly as
well as the presence of inter-chromosomal effects during meiotic recombination [6].

Chromosome Aneuploidy

Down’s syndrome is a complex genetic disease resulting from the presence and expression of 3 copies of the genes located on chromosome 21 (trisomy 21). In most cases, the extra chromosome stems from the failure of normal chromosomal segregation during meiosis (meiotic non-disjunction). The non-disjunction event is maternal in <95% of cases, occurring primarily during meiosis I in the maturing oocyte, before conception. Down’s syndrome occurs with an estimated frequency of 1 in 600 live births and 1 in 150 conceptions [7,8].

Aneuploidy of the X Chromosome

Klinefelter’s syndrome is a form of hypergonadotropic hypogonadism and infertility resulting from a supernumerary X chromosome (47, XXX). Classically, Klinefelter’s syndrome is outlined by gynaecomastia, small, firm testes with hyalinization of seminiferous tubules, hypergonadotropic hypogonadism and azoospermia. The 47, XXX karyotype with different prevalence rate in the general population (0.1%), among infertile patients of azoospermic (11%) and of oligozoospermic men (0.7%) has also been reported. Men having Klinefelter’s syndrome with chromosomal mosaicism (46, XY/47, XXY) are fertile and with non-mosaic or complete, are azoospermic and only a few have any spermatogenesis [6].

Aneuploidy of the Y Chromosome

In 47, XYY syndrome, men are otherwise healthy, while semen analyses frequently indicate oligozoospermia or azoospermia. It has been shown that germ cells with an extra Y chromosome from men with the 47, XXY karyotype have abnormal meiotic pairing, suggesting disrupted meiosis, eventual sperm apoptosis and subsequent oligozoospermia and infertility [9].

Testicular Disorder of Sex Development (DSD)

The DSD also known as 46, XX male syndrome, in which patients have an X;Y translocation with Y-linked gene SRY is placed on one of the X chromosomes. 46, XX males with SRY and testicular DSD have normal male genitalia but show spermatogenesis arrest and develop severe testicular atrophy and azoospermia. The SRY encodes the critical testis-determining transcription factor that activates a number of downstream transcription factors involved in testes formation. SOX9 is a direct target of SRY, and it’s over expression can mimic male development without SRY. Mutations and small duplications of the SOX9 upstream regulatory region were demonstrated in SRY-negative XX males. Alternatively, increased expression of SOX9 can be induced by steroidogenic factor 1, NR5A1 and SOX3. Recently, R-Spondin 1 (RSPO1) mutations were shown to cause an XX male condition [10].

Small Supernumerary Marker Chromosomes

Small Supernumerary Marker Chromosomes (sSMC) are structurally abnormal chromosomes that cannot be identified or characterized unambiguously by conventional banding cytogenetics alone. They can lead both to fertility problems and repeated abortions. The rate of sSMC presence in the normal population was recently determined to be 0.044%, however, elevated to 0.125% in infertile groups. It was identified that after sSMC detection in connection with unexplained infertility in ~60% of cases the origin of the sSMC can be characterized by application of the centromere-specific probes for chromosomes 14 and 15 [11-14].

Myotonic Dystrophy 1

Myotonic Dystrophy 1 (DM1) is a hereditary, autosomal dominant multi-system disorder characterized by the development of structural and functional abnormalities in the muscle membrane protein associated with muscular dystrophy, cardiac conduction disorders, cataracts, mental retardation and endocrine and reproductive defects. Progressive testicular atrophy is a prominent feature and occurs with an incidence of approximately 80%. Histological abnormalities include hyalinization, atrophy, fibrosis of seminiferous tubules and reduced sperm numbers. Oligosperma and azoospermia are also reported in approximately 73% of DM1 patients [15-17].

Single-Gene Disorders

Single-gene mutations are involved in infertility, either by causing aberrant pubertal development, deficiency of pituitary hormones or affecting the gonadal functions. Mutations of genes expressed in the hypothalamus generally result in hypogonadotropic hypogonadism, a condition of absent or deficient puberty owing to low serum gonadotrophin, Follicle Stimulating Hormone (FSH) and Luteinising Hormone (LH). The KAL1 gene is localized in the pseudoautosomal region of the Xp. Mutations like deletions and point mutations cause Kallmann’s syndrome in males. It is an X linked recessive idiopathic condition, associated with hypogonadism and anosmia. Mutations in AHC gene is implicated in adrenal hypoplasia which causes delayed puberty and cryptorchidism in males. Leptin (LEP) mutations posed irreversible pubertal delay [18,19]. Males with FSHβ mutations present with azoospermia, but puberty may be normal or absent in them [20]. SOX9 is a member of a family of transcription factors that contain a Sex determining Region of Y chromosome (SRY) - related HMG box (SOX). Mutations in SOX9 gene have been found in individuals who are chromosomally male but phenotypically female. Still, there are many genes need to be explored in terms of infertility [21,22].

Cystic Fibrosis

The men with Cystic Fibrosis (CF) have been associated with Congenital Bilateral Absence of the Vas Deferens (CBAVD) as a result of which spermatooza are not transported to the urethra, a condition referred to as obstructive azoospermia. Mutations in the CFTR gene have also been identified in patients with CBAVD, which suggests that this condition is a primarily genital form of cystic fibrosis [23-25].

Leydig Cell Hypoplasia

Leydig cell hypoplasia is a rare autosomal recessive condition wherein the fetal Leydig cells are unresponsive to Human Chorionic Gonadotropin (hCG), The condition featured with hypoplasia of the Leydig cells, complete feminization of the external genitals and...
partial masculinization with microepis. Leydig cell hypoplasia is caused by inactivating mutations in the LHCGR gene [26].

**XY Gonadal Dysgenesis**

Gonadal Dysgenesis (GD) can be classified as complete or partial. The gonads in partial GD may be marked by the presence of few tubular structures or fibrous tissues or may occur as streaks. In complete GD, male have a completely female phenotype with no gonadal development. However, complete GD has a higher risk for developing gonadoblastoma [27].

**Female Infertility**

**Advanced Maternal Age and Aneuploidy**

Advanced maternal age has been commonly associated with aneuploidy due to non-disjunction of chromosomes during meiosis. It is considered as main cause of embryonic loss and poor fertility. The "limited oocyte pool" means the lower number of antral follicles in older women's ovaries may cause the recruitment of suboptimal ova. With aging, distribution of chiasmata formed during early prophase I as well as weakened centromeric cohesion; establish a strong predisposition for aneuploidy [6]. 47, XXX syndrome, also known as trisomy X, is one of the most common causes of Premature Ovarian Insufficiency (POI). While, the majority of women with trisomy X present as normal, some suffer from POI or from malformations of the gonitouinary tract [28].

**Turner’s Syndrome**

Turner's Syndrome (TS) is characterized by a complete or partial absence of one X chromosome. The most frequent chromosome constitution is 45X. A mosaic chromosome complement, the most common being 45X/46XX, 46XXq or 46XXp deletions and a ring X chromosome complement can be identified in the syndrome. Thus, the syndrome might be attributable to a limited amount of genetic material in these chromosomes and is usually due to non-disjunction during meiosis. Most women with TS are infertile due to gonadal dysgenesis and a streak ovary composed of white fibrous stromal tissue containing no ova or follicular derivatives. However, at puberty, mostly those with mosaic karyotypes, have ovaries with a relatively low number of follicles, so that there is spontaneous pubertal development [23,30].

**Polycystic Ovary Syndrome**

Polycystic Ovary Syndrome (PCOS) is a complex and heterogeneous endocrine condition marked by hyperandrogenism, hyperinsulinemia, insulin resistance and chronic anovulation. The elevated insulin levels facilitate secretion of androgens from the ovaries and adrenal glands, leading to hyperandrogenism. Elevated levels of androgens lead to menstrual disturbances and infertility. As DNA methylation regulates gene transcription, forty genes have been shown to be differentially methylated in PCOS patients compared with the corresponding genes in normal individuals. Changes in methylation of EPHX1, LMNA, and GSK3A are associated with PCOS. Although, several genes have been associated with PCOS, there is no evidence to suggest that a unique gene or a dominant pathway is the sole causative factor [6,31,32].

**Cystic Fibrosis**

Cystic Fibrosis (CF) is the most common life-shortening genetic disease caused by mutations in the gene encoding a cAMP-regulated chloride channel, the CF Transmembrane Conductance Regulator (CFTR). CF is a systemic illness that affects various organ systems including the pulmonary, endocrine, epithelial, gastrointestinal, pancreatic, immune and reproductive systems. Reduced fertility has also been observed in women with CF. The prominent hypothesis for the decreased fertility in CF females is viscous mucus in the cervix that may create a barrier to sperm passage. Additionally, CFTR is involved in secretion of endometrial and oviduct HCO3-, which is necessary for sperm capacitation. CFTR is also expressed in the cervix, oviduct, ovary and uterus, where it regulates fluid control in the female reproductive tract. CF is associated with menstrual irregularities, including amenorrhea, irregular cycles and anovulation [24,25,33].

**Single Gene Mutations**

FSHβ gene mutation has been shown to cause absent or incomplete breast development, low FSH and oestradiol, high LH and sterility in females [20]. Similarly, Xp11 gene deletions result in ovarian failure as well as affect menstrual function in women [19]. Fragile X syndrome is characterized by mental retardation, long faces, large ears and prominent jaws. The syndrome was first reported in 1969 with constriction of the long arm on the X chromosome. The critical gene for fragile X is FMR1 [34]. Many women with galactosemia manifest hypergonadotropic hypogonadism, presenting with secondary amenorrhea and premature ovarian failure. A candidate gene associated with galactosemia and endometriosis is the GALT gene [35]. Leiomyomas or fibroids are benign tumors found in the smooth muscle layers of the uterus. New studies using conventional and next-generation sequencing techniques identified mutations in the MED12 gene as a major contributor to leiomyoma [36]. Endometriosis is a complex disease, characterized by the inflammation and bleeding of the endometrium. It posed infertility and pain due to endometrial tissue in the pelvic region outside of the uterus. The candidate genes have been identified by genetic association and linkage studies are SNPs and CNVs. Among two Genome Wide Association (GWAS) studies conducted in Australia and Japan in 2011 and 2010, only one common locus was found in the 1p36 region which contains WNT4, a gene responsible for cell proliferation and which plays a key role in embryogenesis involvement in endometriosis [3].

**XX Gonadal Dysgenesis**

Sex determination is controlled by complex molecular signaling and abnormalities in these signaling pathways can lead to gonadal dysgenesis. A well-known illustration of this type is XX female Gonadal Dysgenesis (XX-GD) which is genetically heterogeneous, but phenotypically identified by the presence of gonadal streaks, lack of spontaneous pubertal development, primary amenorrhea, uterine hypoplasia and hypergonadotropic hypogonadism. Ovarian insufficiency can range from lack of pubertal development to the onset of menopause before the age of 40 years. Mutations in FSHR, BMP15, NR5A1, EIF2B2, EIF2B5, HSOD7B4, and HARS2 have been reported in XX-GD [37]. Ovarioleukodystrophy and
Perrault syndrome are examples of syndromic cases of XX-GD. Perrault syndrome is characterized by ovarian dysgenesis, sensorineural deafness, mental retardation, ataxia and cerebellar hypoplasy. Compound mutations at highly conserved amino acids in mitochondrial Histidyl tRNA Synthetase (HARS2) also cause this syndrome. These mutations implicate a role for the mitochondria in proper function of the ovaries [38].

**Premature Ovarian Failure**

Premature Ovarian Failure (POF) is defined as the onset of menopause in women under the age of 40 years. The symptoms include amenorrhea due to hypoestrogenism, elevated gonadotrophin levels and other menopause-related symptoms such as hot flushes, night sweats and vaginal dryness. POF is likely due to depletion of the follicles which could be because of a decreased number of oocytes being formed during development or an increased rate of oocyte atresia during the reproductive lifespan [39]. Paradoxically, the etiology of this disease is not clear and it is likely that the disease is caused by several factors. POF can be influenced by environmental and genetic factors. Irreversible damage to the ovaries during radiation therapy, chemotherapy or autoimmune disease conditions can cause POF. The X chromosome abnormalities and autosomal genetic defects can also cause POF [40]. A number of genes have been associated with POF include FMR1 and Bone Morphogenetic Protein 15 (BMP15). Among autosomal gene mutations often found in women with POF are AR, CDKN1B, CYP19A1, GDP9, FIGLA, FOXL2, FOXO1a, FOXO3a, INHA, LHXB, NOBOX, NANOS3, FSHR and SALL4 [41].

**Conclusion**

The genomic basis of infertility is very complex and is determined by many factors. These factors influence the development of gametes, reproductive organs, their physiology and the development of embryo and its further differentiation. The genetic disorders can affect males, females or both, causing infertility. Genetic disorders can be chromosomal, single gene mutations or can be multi-factorial. Extensive research has been conducted for having a better insight into the genomic basis of infertility. However, inspite of extensive research, there are no well-defined genes that can be used for genetic testing of infertility conditions. Thus, there is a need for newer diagnostic technologies to identify both new and known infertility genes. With the growing incidence of infertility and growing awareness of general population towards newer approach in the treatment of infertility, better understanding in the genetic control of infertility will help in planning treatment modality that would prove beneficial to the infertile couples.

**References**


