Brothers No Longer Sisters: Case Study of Pseudohermaphrodites

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A study was conducted on male pseudohermaphrodites who got transformed from females to males during the years 1993-95, at the age of eighteen and thirty-two. The younger sister (18 years old) transformed before marriage, whereas, the elder sister (32 years old) remained a housewife for about 20 years. No one else in the family was recorded with similar genital anomaly except one ambiguous female. The clinical reports, morpho-anatomical features and chromosomal pattern reveal their testicular feminization/male pseudohermaphroditism.

Keywords: Male pseudohermaphroditism, testicular feminization syndrome, type II 5 α-reductase, androgen receptors

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Introduction
Sex of an individual is determined by genetic events that direct the bipotential fetal gonads either to develop testes or ovaries in an individual (Eicher, 1988). In human beings the presence of Y-chromosome (Page, 1985) determines whether a particular human embryo will differentiate into a male (XY) or a female (XX) baby. However, sometimes the individual determined as male or female develops ovarian or testicular tissues giving rise to the complete sex reversal (Eicher, 1988) or even to pseudosertoli syndromes (Szynder and Davison, 1987). The pseudosertoli syndromes bearing testes with the external genitalia like that of females are referred to as male pseudosertoli syndromes, whereas female pseudosertoli syndromes bear ovaries with the external genitalia like that of males (Thompson and Thompson, 1986). The present study was conducted on male pseudosertoli syndromes where an engaged girl and a married woman got transformed into males at the age of eighteen and thirty-two years respectively.

Materials and Methods
Two individuals, morphologically females, born of the same parents, transformed into males in District Swat, were investigated in this study. The family pedigree was constructed by interviewing the elders of the family and all information was checked by interviewing different persons of the community. The blood samples were collected and chromosomal analysis was carried out according to the method described by Moorhead et al. (1960).

Clinical Report: The 15 years old younger sister (VII-20), an engaged girl, consulted the doctor on complaint of continuous pain in her lower abdomen. She was recognized as male by the doctor and was operated upon followed by gonadotrophin therapy. A year after the transformation, his 32 years old sister, a married woman, was operated upon and transformed into male. Both were females morphologically before transformation, having well-developed mammary glands (gynaecomastia), female external genitalia and large clitorises (clitoromegaly). However, they possessed hirsutism i.e. growth of facial hair and male distribution of pubic hair. They had no menstruation (primary amenorrhoea). The pitch of their voices was initially high but gradually changed towards that of males. According to surgical report, right testes was found in the neck of scrotum and the left one in the scrotum. The scrotum had invaginated in the form of labia majora (labioscrotal fold).

There was no uterus, oviducts or ovaries. The penis was in the form of large clitoris with perineo-scrotal hypospadias. The urogenital slit was closed, the scrotal pouch was corrected and the testes were descended into scrotum. The perineo-scrotal hypospadias was corrected.

Results and Discussion
Case history (Table 1), clinical reports, morpho-anatomical features and chromosomal pattern (44, XY) of the subjects reflect that they were male pseudosertoli syndromes/testicular feminization syndromes as described by Wu et al. 1997. Nordenskjold and Ivansson, 1998. According to Griffith et al. (1993) the frequency of testicular feminization syndrome is one in 65,000 male births. The appearance of female secondary sex characters in these patients at puberty is due to estrogens secreted by the Leydig cells of their testes (Hamm and Veemott, 1980).

Embryological studies reveal that in the 7th week of gestation, normally each embryo has two pairs of male and female primordial genital ducts i.e. mesonephric or Wolffian ducts and paramesonephric or Mullerian ducts differentiating into male internal genitalia (epididymis and vas deferens) and female internal genitalia (oviducts and uterus), respectively (Gilbert, 1997). The differentiation of external genitalia is dependent upon the urogenital slit. If it disappears, formation of male external genitalia results and its persistence gives rise to differentiation of female external genitalia (Sadler, 1995; Ganong, 1997). Development of genital ducts system and external genitalia occurs under the influence of hormones i.e. Mullerian inhibiting substance (MIS) and testosterone circulating in the fetus. MIS and testosterone are secreted by the Sertoli and Leydig cells, respectively, of the fetal testes. MIS causes regression of the Mullerian ducts, whereas testosterone is responsible for maturation of Wolffian ducts into vas deferens and epididymis.

Differentiation of male external genitalia occurs under the influence of dihydrotestosterone which is produced from testosterone by an enzyme, type II 5-alpha-reductase present in genital tissues and prostate, whereas type I 5-alpha-reductase is present in the skin throughout the body including scalp. These two types of enzymes are encoded by different genes (Ganong, 1997) localized on chromosome 5 and chromosome 8, respectively (Thigpen et al., 1992). Male pseudosertoli syndromes or testicular feminization syndrome, where the tissues of external genitalia develop and differentiate like that of females, is a hereditary character (Thompson and Thompson, 1968) which occurs either due to presence of a mutation in androgen receptors gene present on chromosome X and transmitted from a carrier mother to the offspring (Govan et al., 1988; Griffith et al., 1993) or deficiency of type II 5-alpha-reductase encoded by a recessive gene localized on chromosome 2 (Thigpen et al., 1992) or because of defective embryonic testes, which are unable to secrete MIS due to a mutant gene present on chromosome 19 (Cohen-Haguenauer et al., 1987). This condition occurs due to congenital deficiency of 17 alpha-hydroxylase, encoded by a recessive gene present on chromosome 10 (Sparks et al., 1991).

As it is clear from the pedigree (Fig. 1), the family initially comprised of eleven persons i.e. two parents, eight sisters and a brother. Now they are three brothers and six sisters. One of their sisters (VII-10), though had married, could not produce any child and is ambiguous sexually. The family refuses to provide information about her due to tribal norms. Remaining five sisters (VII-6, VII-8, VII-18, VII-21 and VII-22) have married and have children.

Personal history of the subjects shows that they had feminine body contour, and were involved in household activities before transformation. The younger one was engaged and elder one remained a housewife for about 20 years. They were more powerful among their age group females. Emotionally, they behaved as normal females but later on their sentiments inclined towards that of males, however they were regarded as females and they used to present women society. Before transformation, the elder one used to shave his beard and hide his face to other members of the community. After operation, their behavior and emotions were changed so rapidly that they married in a short period of 1-3 years. Presently, the younger one (VII-20) is a father of two children (VIII-9 and VIII-10). His fertility is perhaps due to the fact that previously, his testes were not exposed to the abdominal temperature for longer time as he was diagnosed in the early days of puberty. The elder one has no children and his sterility is perhaps due to the exposure of his testes to higher abdominal temperature for longer time. Chromosomal analysis show a male 44, XY karyotype.

Pinsky et al. (1978) Nordenskjold and Ivansson (1998) reflect that the cases described here are due to the presence of
mutations either in type II 5α-reductase or dihydrotestosterone receptors genes, localized on human chromosomes 2 and X, respectively. If it had been due to mullerian inhibiting substance (MIS) deficiency, the mullerian ducts system would have developed into female internal genitals in the subjects (Behringer et al., 1994; Mishina et al., 1995). Similarly, if the cases had been due to congenital deficiency of 17α-hydroxylase, synthesis of androgens would have not taken place resulting in neither differentiation of male internal genitals (Walter et al., 1998) nor feminization in the subjects.

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