Transverse Testicular Ectopia: A Rare Case Report

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Abstract:
Transverse testicular ectopia (TTE) is a rare but well-known congenital anomaly that occurs 1 in 4 million in which both testes migrate toward the same hemiscrotum. In most of the cases it is an intra-operative finding, but preoperative diagnosis can be made by careful history taking, physical examination and imaging studies. Further evaluation is very important because it can be associated with other congenital anomalies. We report a case of TTE in a 32 years old male who presented with sudden painful swelling in right inguinoscrotal region. Physical examination revealed right sided obstructed inguinal hernia and left sided non palpable testis with underdevelopment of left hemiscrotum. On exploration, one testis is found within the hernial sac and the other testis within scrotum of same side. The testis which was already in right side of scrotum was kept in same place and the other testis which was found within hernial sac was kept in subdartos pouch at the root of right side of scrotum.

Key words: Transverse testicular ectopia, Inguinal hernia.

Introduction:
The first known case of transverse testicular ectopia (TTE) was reported by Von Lenhorsek in 1886 of a 35-year-old adult¹. In the literature more than 100 cases of TTE have been reported². Other anomalies like persistent Mullerian duct syndrome, renal agenesis, hypospadias, hermaphroditism, and scrotal anomaly can be associated³,⁴. An inguinal hernia is commonly present on the side to which the ectopic testis has migrated⁵. The ectopic testis may lie in the opposite hemiscrotum, in the inguinal canal or at the deep inguinal ring. The main treatment of TTE is surgical intervention with hernia repair, if present⁶.

Case Report:
A 32 years old male patient presented to emergency department with sudden painful swelling in right inguinoscrotal region, colicky abdominal pain and vomiting for 4 hours. He gave history of right sided inguinoscrotal swelling for last 4 years which appeared on standing and coughing and disappeared on lying down with occasional dragging pain. Physical examination revealed right sided obstructed inguinal hernia and left sided non palpable testis with underdevelopment of left hemiscrotum. He is father of two children. With diagnosis of right sided obstructed inguinal hernia and left sided undescended testis, he underwent emergency operative treatment under general anaesthesia after adequate resuscitation. Right oblique inguinal incision was given and after incising external oblique aponeurosis the cord structures were identified. Hernia sac was opened which contained omentum and small gut which were initially little blackish in colour but returned to normal pink colour after incising the constricting band at deep inguinal ring; were reduced into peritoneal cavity. One unusual finding was discovered at this stage: one testis was found within hernial sac along with the normally situated testis within scrotum of same side. Both testicles were delivered into the inguinal wound (Figure 1). Both spermatic cords were followed to right deep inguinal ring where they entered the abdomen. Isolation of hernial sac was carefully done in regular way without any injury to the structures of spermatic cord. The testis that was found within hernial sac was kept in subdartos pouch at the root of right side of scrotum and the testis that was already in right side of scrotum was kept in same place without any orchidopexy. So right side of scrotum containing two testes; one in original position and another at subdartos pouch at root of right scrotum (Figure 2). Postoperative period was uneventful and follow up after 8 weeks shows positioning of two testes in right side of scrotum; one at the bottom of right side of scrotum and one at the root of right side of scrotum with empty and under development of left side of scrotum. The patient had no complaints.

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Discussion:

TTE is a rare form of testicular ectopia. Many different nomenclatures such as crossed testicular ectopia, testicular pseudo duplication, unilateral double testes and transverse aberrant testicular maldescent have been used for this condition. Most cases of TTE described in the literature are diagnosed before the age of 18 and management is targeted to protect fertility and reconstruct a normal anatomy by transferring the testis and repairing any associated anomalies such as inguinal hernia. One case of family incidence was reported by Staubes, in two brothers with TTE and persistent Mullerian remnant. The patient generally presents with an inguinal hernia on one side and a contralateral cryptorchidism. The precise etiology of TTE is not clear yet. Few theories exist regarding the embryogenesis of TTE viz. adhesion and fusion of developing Wolffian ducts, aberrant gubernaculum, testicular adhesions, defective formation of the internal inguinal ring, and traction on a testis by persistent Mullerian structures. Berg described the possibility of the development of both testes from the same genital ridge. Gupta and Das suggested that due to early adherence and fusion of the developing Wolffian ducts, the descent of one testis has caused the second one to follow.

There are five known types of testicular ectopia: superficial inguinal (interstitial), femoral (crural), perineal, pubopenile, and crossed (transverse). According to Gauderer et al, three types of transverse ectopia are recognized: Type I, which is associated with an inguinal hernia and accounts for 40-50% of the cases; Type II, which is accompanied by Mullerian duct remnants (30%); and type III (13-20%), which includes genitourinary anomalies other than persistent Mullerian duct such as hypospadias, pseudo-hermaphroditism, bifid scrotum, renal anomalies, seminal vesicle contralateral aplasia, and seminal vesicle cysts. According to that classification, our case was type 1 TTE.

Patients with TTE are at increased risk of malignant transformation. In fact, the overall incidence of malignant transformation in these testes is 18%, similar to the rate in abdominal testes in otherwise normal men. There are reports of embryonal carcinoma, seminoma, yolk sac tumor, teratoma and spindle cell neoplasm- leiomyoma with TTE.

For preoperative diagnosis, along with high suspicion the imaging investigations include inguinoscrotal ultrasound and transabdominal ultrasound, MRI of abdomen and pelvis and contrast-enhanced CT is needed to look for associated anomalies.

Once diagnosis of TTE is made, a conservative surgical approach in the form of orchiopexy is recommended for the preservation of fertility. Laparoscopy is useful for both diagnosis and treatment of TTE and associated anomalies. Management for TTE is either trans-septal or extra-peritoneal transposition orchiopexy along with a search for other anomalies. Long-term postoperative follow-up is recommended.
**Conclusion:**

TTE is a rare but well studied congenital anomaly. The ectopic testis can lie in the hemiscrotum, in the inguinal canal, or at the deep inguinal ring. The diagnosis should be suspected in patients presenting with inguinal hernia on one side and cryptorchidism on the other side. In suspected cases, ultrasonographic evaluation is very much helpful in diagnosing this condition before surgery. Transseptal orchiopexy is the recommended surgery to perform. At present, laparoscopy is useful for both diagnosis and management of TTE and associated anomalies.

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