TRANSVERSE TESTICULAR ECTOPIA

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ABSTRACT… Crossed testicular ectopia (CTE) is a rare congenital anomaly¹⁴. It is also known as transverse testicular ectopia (TTE). In this condition both testis descend/migrate towards the same hemi-scrotum. About 100 cases have been reported so far. It may be associated with other congenital anomalies which include persistent Mullerian Duct Syndrome, true hermaphroditism, pseudo-hermaphroditism, inguinal hernia, hypospadias and scrotal anomalies. We are reporting one such case in 6 years old boy who presented to CMH Malir in March 2014 with left inguinal hernia and per-operatively found to have right testis in left inguinal canal. Left testis had descended into left upper scrotum. Vasa were tethered together in upper portion by fibrous tissue. Right testis and vas was separated, mobilized extra-peritonially and done in right scrotum (ipsilateral). Left orchiopexy was done in left scrotum. Hernia repair was then preformed. Post-op u/s scan was done to detect any other associated pathology which was negative. In most of the published reports diagnosis was made per-operatively as was the case in this patient.

Key words: Testis, Undescended testis, Cryptorchidism, Ectopic, Hemi-scrotum, TTE

INTRODUCTION

Transverse testicular ectopia (TTE) is a rare condition, (about 100 cases reported so far)¹. It is also named crossed testicular ectopia, testicular pseudo-duplication, unilateral double testis and transverse aberrant testicular maldescent. In this condition both testis descend through the same inguinal canal towards the same hemi-scrotum. In published reports it is usually the right testis which traverses the left inguinal canal. However reports are also there in which left testis had descended towards right hemi-scrotum. Pre-operative localization on ultrasound by an experienced sinologist may be helpful but diagnosis is usually made during surgical exploration for hernia repair.

CASE PRESENTATION

A 06 year old boy presented to Combined Military Hospital (CMH Malir) in Mar 2015 with a lump in left inguinal region which increased in size on straining. It was reducible and cough impulse was present. Left testis was palpable in left upper scrotum. Right testis was not palpable. Diagnosis of left inguinal hernia with undescended right testis was made. Birth history was normal spontaneous vaginal delivery at home. Herniorrhaphy planned as first stage as this was the presenting complaint. During operation another well-developed testis was found in left inguinal canal close to the internal ring in addition to the left testis which was descended in left upper scrotum. Vigilant dissection revealed that its blood supply and vas were coming from right side. The two vas were adherent with fibrous tissue but clearly separate. Two vas were separated and right testis was mobilized extraperitoneally around the root of the cord and done in right scrotum. Left orchiopexy was done in left scrotum. Hernia repair was then performed. Post-op u/s scan was done to detect any other associated pathology which was negative. In most of the published reports diagnosis was made per-operatively as was the case in this patient.

Figure-1. Both testes seen in the left hemi-scrotum
penis. Right orchiopexy was done in ipsilateral right hemi-scrotum. Left orchiopexy was done too and hernial defect was repaired. Post-op ultrasound of abdomen and pelvis did not reveal any abnormality.

DISCUSSION

TTE is a rare clinical entity. Usual sites of ectopic testes are perineum, thigh, and anterior abdominal wall. Opposite inguinal canal and scrotum are the rare sites of ectopia. The testis migrates to the opposite side inguinal canal instead of the same side. It was first described by Van Len Hossek in 1866. Several theories have been reported to explain the genesis of TTE. Berg proposed the possibility of development of both testes from the same genital ridge. Kimura found that if both vas deferens arise from one side, there had been unilateral origin but if there was bilateral origins, one testes had crossed over. Gupta and Das postulated that adherence and fusion of the developing Wolffian duct might have occurred which resulted in the descent of both testis to the same side.

Another theory is that descending testis instead of following the main gubernaculum during descend to the ipsilateral hemi-scrotum, follows one of the additional branches of gubernaculum and ends up at ectopic site. One such branch goes to the opposite inguinal canal and contralateral hemi-scrotum. Following this branch will result in TTE. Inguinal hernia is almost invariably present on the side to which the ectopic testis has migrated.

TTE has been classified in to three types:
Type I Accompanied only by hernia (40-50%).
Type II Accompanied by persistent or rudimentary Mullerian duct structures (30%).
Type III Associated with anomalies like hypospadias, pseudohermaphroditism and scrotal anomalies (20%), other urinary tract anomalies.

According to this classification, our case was of type I TTE. Association with fused vas deferens is extremely rare. This condition obstructs the ipsilateral orchiopexy during surgery. Mean age of presentation is 4 years. Few cases have been reported among adults (late diagnosis) Clinical presentation is usually with inguinal hernia with contralateral or bilateral cryptorchidism. Contralateral undescended testis is typical but usually missed because of the rarity of this condition. Diagnosis is made generally at surgical exploration as was there in our case. Pre-operative ultrasound by an experienced sonologist might help. Laparoscopy maybe helpful in both diagnosis and treatment of TTE. Patients with TTE are at increased risk of germ cell testicular tumors (18%). Embryonal carcinoma has also been reported. Walsh et al concluded that the chances of malignancy are six times higher in cryptorchidism where operation was delayed until after 10 years. After mobilization of cord the ectopic testis can be fixed in the ipsilateral hemi-scrotum (orchiopexy) either extra-peritoneal or trans-septal. In cases where vas deference’s are fused and it is difficult to perform extra-peritoneal ipsilateral orchiopexy Trans-septal orchiopexy can be considered.

CONCLUSION

TTE is a rare congenital anomaly. Exact pathogenesis is not clear. Fusion of Mullerian duct or descent along a rudimentary branch of gubernaculum are possible. Testis may be present in opposite hemi-scrotum, inguinal canal or at deep ring. Diagnosis should be suspected when a child presents with unilateral inguinal hernia and opposite testis is not palpable. However, diagnosis is usually made during surgery. Pre-operative ultrasound and laparoscopy may help in diagnosis if suspected clinically. Extra-peritoneal ipsilateral orchiopexy is recommended when vas and its vessels can be mobilized easily. In difficult cases of vas fusion, trans-septal orchiopexy should be considered. Post-operative scan should be done for any rudimentary uterus (Mullerian duct syndrome), pseudo-hermaphroditism.

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REFERENCES


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