Triorchidism: presenting as undescended testis in a case of indirect inguinal hernia

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Abstract

Triorchidism is the commonest variety of polyorchidism, an entity with more than two testes. It is an extremely rare congenital anomaly of the testis. Although excision of the abnormal testis is a safer alternative proposed, recent literature suggests more conservative approach in normal testes with watchful regular follow up to screen for malignancy. This case presented as a left inguinal swelling diagnosed as indirect left inguinal hernia. The left side testis was of smaller size (about half) with normal sperm count, morphology and motility. Intraoperatively indirect inguinal hernia was noted with supernumerary testis at deep ring in addition to normal left testis in left scrotal sac. The ectopic testes were small (2.5x2.5x1 cm) lacking epididymis and with short vas deferens. An evident normal semen analysis showed normal sperm count, morphology and motility. The patient was operated for left inguinal hernioplasty. Intraoperative findings of indirect inguinal hernia sac was present with small undescended super-numerary testis of size 2.5x2.5x1 cm at the deep inguinal ring (Figure 1) having short vas deferens and a normally located left testis with epididymis and vas deferens in the scrotal sac. Due to anomalous anatomy of super-numerary inguinal testis, orchidectomy with excision of indirect sac was performed. The normally located left testis in scrotal sac was preserved. Patient recovered uneventfully. The histopathological examination of the specimen reported thickened capsule of testis and tubules with arrest of development of the germ cells (Figure 2). A prominence of Sertoli and Leydig cells along with hyalinised, thickened basement membrane of seminiferous tubules and dense intervening stroma was found. A follow up tumor marker assay showed negative study, which can likely be present in burnout tumor in spite of negative histology.

Introduction

Polyorchidism is the incidence of more than two testes. There is an extremely rare congenital anomaly, but well documented in literature. The first case was been reported in 1880. There are about 190 cases reported with higher prevalence of diagnosis in pediatric age group. With first histological confirmation in 1895 there are about 140 histologically proven cases. Polyorchidism can clinically have various presentations as inguinal swelling, cryptorchidism, testicular torsion, hydrocele and varicocele with majority being asymptomatic. The most common form of polyorchidism is triorchidism (or 3 testes) with left sided predominance (63%) over the right. A higher tendency for malignant transformation in non-functional testis necessitates the radiological and functional assessment of polyorchid testes. We hereby report a case of 24-year old male with left side super-numerary non-functional testis, presenting as left sided inguinal hernia.

Case Report

A 24-year old young male came with reducible swelling in left inguinal region since 4 years without associated history of trauma or pain. On detailed local examination, a reducible swelling in the left inguinal region with positive cough impulse and positive deep ring occlusion test was noted. The right testis was normal in size, whereas left testis was smaller in size 3x3x2 cm (about 50%) compared to the right side. The vas deferens on both sides was distinct and normally palpable. Sensation over testis was normal without any significant inguinal lymphadenopathy.

The patient was investigated for surgical fitness. Semen analysis showed normal sperm count, morphology and motility. The patient was operated for left inguinal hernioplasty. Intraoperative findings of indirect inguinal hernia sac was present with small undescended super-numerary testis of size 2.5x2.5x1 cm at the deep inguinal ring (Figure 1) having short vas deferens and a normally located left testis with epididymis and vas deferens in the scrotal sac. Due to anomalous anatomy of super-numerary inguinal testis, orchidectomy with excision of indirect sac was performed. The normally located left testis in scrotal sac was preserved. Patient recovered uneventfully. The histopathological examination of the specimen reported thickened capsule of testis and tubules with arrest of development of the germ cells (Figure 2). A prominence of Sertoli and Leydig cells along with hyalinised, thickened basement membrane of seminiferous tubules and dense intervening stroma was found. A follow up tumor marker assay showed negative study, which can likely be present in burnout tumor in spite of negative histology.

Discussion

Polyorchidism presents itself more commonly in the 2nd to 3rd decade, with approximately 50% of cases reported between 15 to 25 years of age. The exact etiology is unknown. The common presentations of polyorchidism manifests as maldescend (40%), hernia (30%), torsion (15%), hydrocele (9%), malignancy (6%), infertility and epididymitis.

Table 1 describes the Bergholz classification of polyorchidism. The testis takes its origin from the medial part of primitive genital ridged while the epididymis and vas deferens develop from the Wolffian duct. The exact etiology of polyorchidism is unclear, however accidental division of genital ridged before 8 weeks of gestation could be a possible cause. There are theories proposing the probable presence of multiple testes first being duplication of longitudinal genital ridged resulting into separate testes so that total volume of testis exceeds of one. The second theory explains transverse division of genital ridged at different levels resulting into different combination of testes, epididymis and vas deferens.

There are various types of classifications for polyorchidism. Based on site of testes they can be classified as scrotal (66%), inguinal (23%) and abdominal (9%). Leung (1988) classified polyorchidism based on embryological development with possible anatomical distinct epididymis or vas deferens. Singer (1992) and colleagues proposed classification based on anatomical and functional potential of supernumerary testis to epididymis and finally by vas deferens.

An effort to restore the functionality of undescended testes can be attempted if surgically corrected before age of 4-5 years. A significant risk of malignancy is reported in undescended non-functional testes before puberty being 2.2% while after puberty it rises substantially to 5.4%. Recent publications also suggest higher incidence of testicular malignancies (4 to 6%) in cryptorchid testis compared to normal testis. There is higher risk of testicular malignancy in polyorchidism cases of about 7% especially in non-scrotal testis.
these 88.8% are malignant namely seminoma, teratoma, yolk sac tumors and choriocarcinoma.\textsuperscript{4} About 20% of polyorchidism patients seek the help of doctors for infertility issues.\textsuperscript{2}

The management of supernumerary testes has been a topic of much debate and different factors like age of patient, location of testes, functional status and lastly compliance and accessibility for follow up have to be taken into consideration. Broadly in pediatric age group or patients with reproductive desire and uncomplicated polyorchidism should have conservative approach like orchidopexy,\textsuperscript{8} however a strict vigilant follow up is mandatory. While adults without reproductive desire, complicated polyorchidism and with inaccessibility for follow up should not be considered for conservative management.

In our view supernumerary testis with complications like cryptorchidism, torsion and intraoperative non-functional testis i.e., without a draining vas deferens or epididymis,\textsuperscript{7,8} excision can be performed due to higher risk of malignancy in polyorchidism.\textsuperscript{7}

### Conclusions

The diagnosis of uncomplicated polyorchidism is rarely non-incidental and management being controversial.

A surgical approach is advisable in complicated, non-compliant and non-reproductive patients in view of higher risk of malignancy in these patients.

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