Introduction

Polyorchidism, or supernumerary testes is a rare condition, which is embryological in origin. It results from an abnormal division of the genital ridge. Although three testes is the most common form, as many as five have been reported. In approximately 75% of cases, the supernumerary testes are intrascrotal, and the patients most often present with a painless scrotal mass. Of the remaining cases, 20% of the testes are inguinal and 5% are retroperitoneal.[1,2] Supernumerary testes are retractile and more mobility is at increased risk for torsion. An increased prevalence of carcinoma has also been reported.[1]

Case Report

A 72-year-old man presented with vague pain in the lower abdomen and scrotum. There was no history of trauma and he stated that the pain was severe in nature and had been present for a short time. He was married and had three children. Physical examination revealed swelling of scrotum on the left side. There were no features to suggest inguinal hernia. He was referred to the radiology department for evaluation of pain.

Ultrasonography of abdomen was normal. Gray scale and color Doppler sonography of right hemiscrotum revealed a normal testicle with homogenous texture, measuring about $3.8 \times 3$ cm. Normal intratesticular flow was noted. Epididymis was normal in size and echotexture. Minimal hydrocele was noted [Figure 1]. Left hemiscrotum revealed two testicles The lower testis showed homogenous echotexture, measuring $3.7 \times 3$ cm, with normal intratesticular flow. The upper testis measured $3.2 \times 2.4$ cm and showed homogenous hypoechogenicity with no detectable intratesticular flow compatible with ischemia [Figures 2 and 3]. Epididymis was single and appeared normal. Minimal hydrocele was noted.

Magnetic resonance imaging (MRI) examination was done, which showed normal T1 and T2 images of right testis and lower testis of left hemiscrotum [Figure 4]. The upper testis of...
left hemiscrotum showed increased signal intensity on T1 and decreased signal intensity on T2 images, suggestive of torsion and hemorrhagic necrosis [Figure 5]. Minimal hydrocele was noted in both hemiscrota.

Discussion

On the basis of embryologic development, Leung classified polyorchidism into four types[3] In type A, the division separates a small part of the genital ridge, which does not contact the mesonephric duct. Therefore, the supernumerary testis lacks an epididymis and vas deferens. In type B, the division of the genital ridge occurs in the region where the primordial gonads are attached to the mesonephric ducts and the supernumerary testis has its own epididymis. In type C, the supernumerary testis has its own epididymis and shares the vas deferens with the regular testis in a parallel fashion. In this type of polyorchidism, there is an incomplete longitudinal division of the genital ridge and the proximal portion of the mesonephric duct. In type D, which is the least common, complete longitudinal duplication of the genital ridge and mesonephric duct occurs, with resultant complete duplication of testes, epididymides, and vas deferens. Further investigation could not be done in our case to relate as per Leung classification.

Triorchidism is the most common type of polyorchidism and presents with two testes on one side (usually the left) and one testis on the other side. Rare case of polyorchidism with three homolateral testes on the right side and absent testis on the left side has been reported.[4] Splenogonadal fusion is a rare congenital anomaly that may sonographically resemble polyorchidism.[5,6] In this entity spleen, gonad, epididymis, and vas deferens are fused. Sonography reveals a mass with the testicle of similar echogenicity and it may mimic the appearance of polyorchidism. When splenogonadal fusion is suspected, a technetium sulfur colloid scan should be performed to confirm the presence of ectopic splenic tissue.[5] Singer et al.[7] proposed a classification based on reproductive potential of the supernumerary testis. In type 1, the supernumerary testis has reproductive potential because of attachment to a draining epididymis and vas deferens. In type 2, the supernumerary testis has no reproductive potential because of lack of a draining system. Singer et al. further subclassified these types into A, if the supernumerary testis is located in the scrotum, or B, if it is ectopic. The sonographic appearance of polyorchidism is presence of scrotal mass that
has an echo pattern identical to that of the ipsilateral testicle.\textsuperscript{[8]} MRI appearance is a round or oval shaped structure showing typical signal characteristics of testicles, i.e., homogeneous intermediate signal intensity on T1 weighted and high signal intensity on T2 weighted images.\textsuperscript{[9]}

Polyorchidism is a rare genitourinary abnormality, and torsion is one of its associated complications; however, the diagnosis of polyorchidism with or without torsion can be made readily with ultrasonography when one is aware of this entity.\textsuperscript{[10]}

Sonographic features of torsion are homogenous hypoechoegenicity of testis with absent flow on color Doppler study. MRI features of torsion are increased signal intensity on T1 and decreased signal intensity on T2 weighted sequences.\textsuperscript{[11]} Our case showed features of polyorchidism with torsion of one of the testes in left hemiscrotum, both on sonography and MRI.

**Conclusion**

Polyorchidism is a rare congenital anomaly. Torsion is one of the complications of this entity. Polyorchidism can be diagnosed on sonography. Torsion can be suspected and diagnosed on color Doppler study. MRI can be used for confirmation but may be more helpful in cases complicated by torsion, cryptorchidism, or neoplasia.

**References**