Splenogonadal Fusion: A Rare Benign Testicular Mass in a 55-Year-Old Male

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ABSTRACT
Splenogonadal fusion (SGF) is a rare cause of testicular mass with the vast majority of cases presenting in men under the age of 30 and represents a diagnostic challenge. Discontinuous splenogonadal fusion presenting as a new testicular mass in a 55-year-old man is discussed to aid other surgeons in diagnosing this condition.

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Introduction
Splenogonadal fusion (SGF) is a rare congenital anomaly where splenic tissue fuses with gonadal tissue likely during the fifth to eighth week of gestation and is rare with less than 200 cases reported. SGF can be continuous, a fixed connection between the spleen and gonad, or discontinuous, no connection to the spleen. The vast majority of cases are in males under the age of 39 and involve the left gonad. SGF can present in the workup of infertility, cryptorchidism, inguinal hernia, or as a testicular mass as presented in this case.

Case report
A 55-year-old male presented to the urology clinic with obstructive voiding complaints. Past medical history was significant for cardiac disease, otherwise unremarkable. On physical exam he was found to have a 3 cm firm left sided upper pole testicular mass. On further questioning, he had never noticed this mass nor had any of his providers. Scrotal ultrasound revealed a well-circumscribed, focal, solid intratesticular lesion measuring 2.4 × 2.5 × 2.4 cm suspicious for testicular mass (Fig. 1). Serum AFP, HCG, and LDH were negative. Chest X-ray was negative for metastatic disease. CT of the abdomen and pelvis showed no adenopathy in the retroperitoneum. Given the appearance on ultrasound and the history of new testicular mass, left radical inguinal orchiectomy was performed uneventfully. The pathology report revealed a spermatic cord lipoma, benign testicle and epididymis, and splenogonadal fusion with splenic tissue involving the superior pole, 2.5 cm in maximal diameter. The splenic tissue...
had a well delineated fibrous encapsulation, separate from the testicular tissue (Figs. 2 and 3).

Discussion

A new solid intratesticular lesion in a 55-year-old male is considered malignant until proven otherwise. Although, the patient underwent the correct diagnostic workup and treatment given his presentation, he was found to have a rare benign lesion. The question arises as to whether this could have been diagnosed prior to surgery, thus, preventing radical orchiectomy.

The majority of SGF cases are in younger men, continuous, and up to 26% are noted to have other congenital anomalies making the diagnosis more apparent.2 The patient may also give a history of a long standing stable scrotal mass and ultrasound may show a very well circumscribed, rounded, slightly hypoechoic, homogeneous, focal mass. Such findings could prompt the physician to approach the mass in a testis sparing fashion, with intraoperative frozen section to confirm the diagnosis as well as preserve the remaining testis.3 However, testicular tumors have been found concomitantly with SGF, further complicating the diagnosis.2 When diagnostic indicators are absent, SGF represents a significant diagnostic challenge, requiring a high index of suspicion. Given the rarity of SGF, it may be difficult to recommend technetium scans for all patients presenting with testicular lesions. Yet, surgeons can also identify the tumor in the operating room by sending frozen sections if the diagnosis remains in question.3

Conclusion

SGF remains a diagnostic challenge, but remains a consideration in the differential diagnosis of a well circumscribed testicular lesion, especially in men presenting with a long history of a stable testicular mass. Technetium scan and testis sparing surgery with frozen section analysis may help aid in the diagnosis and prevent radical orchiectomy. Nevertheless, a new onset testicular mass should likely be considered malignant until proven otherwise.

Conflicts of interests

None.

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