Neoplastic diseases of the spermatic cord: an overview of pathological features, evaluation, and management

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Abstract: Extracellular tumors found with the spermatic cord, known as neoplasms, are usually identified to be benign. However, the accurate and timely diagnosis of spermatic cord masses is highly crucial, especially when most results are often overlooked or unclear. In this review, we discuss the anatomy and embryology of the spermatic cord. Upon rooting these fundamental concepts, we discuss an array of benign and malignant neoplastic tumors, including their origin, pathological features, clinical evaluation and management, as well as other case-specific characteristics of unique presentation. Many of these neoplasms are based on local neurological, vascular, muscular, bone, soft tissue, or lymphatic origin, while others have metastasized from particular areas of the body.

Keywords: Spermatic cord; spindle cells; myofibroblastic tumor; malignancies; neoplasm; aggressive angiomyxoma; myositis ossificans; scrotum

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Introduction

Neoplasms of the spermatic cord are very rare. These tumors are classified as extracellular and most are benign. They usually originate in the mesothelial layer. Along with benign neoplasms, malignant neoplasms are classified and listed on Table 1.

Diagnosing of the spermatic cord masses is critical during the clinical examination. This region is often overlooked and examination results are unclear. If mass suspicion is detected via palpation during a physical examination, the primary imaging modality used is ultrasonography (US) to visualize the mass. Other imaging modalities should be used to further visualize the mass and prepare for surgery to excise the mass.

Anatomy & embryology

Anatomy

The spermatic cord begins at the inferior abdomen and ends at the scrotum. The contents of the spermatic cord are bound together and covered by three fascial layers: external spermatic fascia, cremaster muscle and fascia, and internal spermatic fascia. All three fascial layers are covered by another layer, known as the superficial fascia, which is found directly below the scrotal skin. The spermatic cord contains many structures, vessels, and nerves which supply blood and signals to and from the testis. The major structure found in the spermatic cord, is the testicular artery, branching off the aorta, inferior to the renal arteries. Next is the artery and vein supplying blood to the cremasteric fascia and muscle, via the
The artery to the vas deferens is also found in the spermatic cord, branching off the inferior vesicle artery, which arises from the internal iliac.

**Embryology**

Formation of the spermatic cord is done at the opening of the inguinal canal, also known as the deep inguinal ring.

**Benign neoplasms**

**Adenomatoid tumor of the spermatic cord**

Adenomatoid tumor of the spermatic cord is a rare neoplasm, which is classified as a paratesticular tumor. These tumors originate in the mesothelium (1). There are very few cases of primary adenomatoid tumor of the spermatic cord in the literature. Miyoshi et al. presented a case of adenomatoid tumor of the spermatic cord. The tumor was asymptomatic and was untreated for years, after the patient had noticed an irregularity in the scrotum. Palpation was used to identify the presence of a mass. The tumor was removed surgically under local anesthesia. Histology of this tumor is very unique, presenting with lumens on the smooth muscles and vacuoles of lymphatic nodules (2).

**Angiomyofibroblastoma of the spermatic cord**

Angiomyofibroblastoma of the spermatic cord is a benign soft tissue tumor. There are only two reported cases of primary angiomyofibroblastoma of the spermatic cord. Tzanakis et al. presented a case of a man with an asymptomatic mass. A surgical procedure was done. The tumor consisted of spindle-shaped cells, uniquely differentiating it aggressive angiomyxoma and other malignant tumors (3).

**Angioleiomyoma of the spermatic cord**

Angioleiomyoma of the spermatic cord is a rare, benign lesion. Ghei et al. presented a case of this rare tumor. Using US and magnetic resonance imaging (MRI), the mass was identified. Scrotal exploration was conducted to learn more about the mass. Local excision was conducted to remove the tumor and is the accepted treatment option (4).

**Angiofibrolipoma of the spermatic cord**

Angiofibrolipoma of the spermatic cord consists of adipocytes, vascular tissue, and collagenous connective tissue, observed through pathological studies. Physical examination identified the presence of a mass in the right spermatic cord. Using US, a cystic mass was found. Under general anesthesia, the tumor was excised from the spermatic cord, with no reoccurrence in the follow-up (5).

**Dermoid cyst of the spermatic cord**

Dermoid cyst, also known as teratoma, arises in the epidermis. It can present itself in children (6) or adults (7). Dermoid cysts are common, but rare originating in the spermatic cord (6). Salemis et al. presented a rare case of a large cyst which presented as an incarcerated inguinal hernia. Lichtenstein polypropylene mesh repair was used to treat dermoid cyst of the spermatic cord (8). Skorniakov et al.
presents a rare case of dermoid cyst of the spermatic cord. In this case, physical examination and US were inconclusive in identifying the mass, so a surgical exploration was conducted and excision was done to remove the mass (7).

**Diverticulum of the spermatic cord**

Diverticulum is an outpouching of a hollow structure in the body. A unique case of diverticulum that involved the spermatic cord was presented by Grigor et al. This case of diverticulum of the spermatic cord was presented as asymptomatic. The discovery was made during a repair of inguinal hernia. A nodule was found similar to a melanoma metastasis attached to the spermatic cord. Upon excision, histology report indicated a detached colonic diverticulum, which parasitized onto the spermatic cord (9). There is no report of primary diverticulum of the spermatic cord in the literature.

**Fibrous pseudotumor of the spermatic cord**

Fibrous pseudotumor of the spermatic cord is classified as a rare benign tumor, which most commonly occurs in the paratesticular region, originating from the intrascrotal tissue. Dieckmann et al. presented two cases of fibrous pseudotumor of the spermatic cord. In both cases, the mass was painless. Original suspicion during the clinical examination was a malignant growth. US confirmed the mass on the spermatic cord in the first patient. MRI was used to confirm a mass in the second patient. Patients underwent local excision to remove the mass. Follow-ups indicated no recurrence in the patients. Histology found collagen-rich, fibrous tissue (10).

**Fibrolipoma of the spermatic cord**

Fibrolipomas are subtype of lipoma, which are benign neoplasms. Fibrolipoma of the spermatic cord is a rare entity. Terada et al. presented a case of an individual with fibrolipoma of the spermatic cord. The patient found a mass in the inguinal region. Tumorectomy and right orchiectomy was used to excise the tumor. Microscopic study showed the tumor composing of adipose tissue (11). Diagnoses can be done using US and computerized tomography (CT), which clearly identifies the mass (12).

**Heterotopic ossification of the spermatic cord**

Heterotopic ossification of the spermatic cord is a rare case of bone formation. The literature has reported very few cases of this pathology. Demirci et al. presented a case of a patient with bone development of the spermatic cord. The patient had a history of pain and swelling and scrotal tenderness. The patient tried self-medication using anti-inflammatory agents to reduce pain and swelling. A physical examination showed a long mass localized to the spermatic cord and laboratory studies were clean. US and MRI were used to diagnose the mass, with both depicting similar findings. The treatment approach for the patient was decided as a radical orchiectomy (13).

**Mesothelial cyst of the spermatic cord**

Mesothelial cyst of the spermatic cord is a rare pathology, and often misdiagnosed as cryptorchidism. Excision of the mass was done. Patient recovered normally without any complications. There was no retraction of the testis into the groin (14).

**Myositis ossificans of the spermatic cord**

Myositis ossificans is the calcification of the muscle, most commonly in the arms or quadriceps. Ozgür et al. presented a case of a patient admitted through outpatient clinical, because the patient noticed a mass growing in the inguinal region. Physical examination revealed a hard mass which was painless, combined with hydrocele. Imaging modalities, US and MRI, were used to locate the mass. Blunt and sharp dissection was used to separate the mass from surrounding structures and orchiectomy to remove the mass from the spermatic cord. Microscopic examination revealed bone structure with dense fibrous tissue (15).

**Neuroectodermal tumor of the spermatic cord**

Primary neuroectodermal tumor of the spermatic cord is a rare neoplastic manifestation of a neoplasm. There are only two cases of this pathology in the literature. In both cases, the tumor was asymptomatic. The mass was palpable, and diagnostic studies such as US, MRI, and CT were used to confirm the mass. Surgical excision of the tumor was performed using radical orchiectomy. Histology and immunohistochemical staining was used in both cases to confirm the diagnosis of neuroectodermal tumor, which presented with small round cells. The immunohistochemical staining was inconclusive for CD99 (16) in one case but not the second case (17), but confirmed for vimentin and CD56 (17). In both cases, chemotherapy was
advised to control the tumor reoccurrence (16,17).

Neurofibroma of the spermatic cord

Neurofibroma of the spermatic cord is a rare pathology. It is found on the peripheral nerve, composed of Schwann cells and fibroblasts. Though these lesions are benign, they have the potential to cause significant morbidity and psychological distress. US, MRI, CT, and positron emission tomography can all be used to identify the presence of the mass. Surgical resection is the best means of treatment and final diagnosis must be made using histopathology (18-21).

Paraganglioma of the spermatic cord

Presentation of paraganglioma of the spermatic cord is asymptomatic. Kwon et al. recently described a case of this pathology. US clearly identified the mass and orchiectomy was used to remove it. Histopathology showed features of malignancy, PET-CT was done on the whole body to determine metastasis of the tumor. PET-CT did not find any other lesions (22).

Spindle cell lipoma of the spermatic cord

Spindle cell lipoma commonly occurs as a solitary subcutaneous, circumscribed lesion in the back. Manifestation on the spermatic cord is an extremely rare occurrence. Ide et al. presented a case of a patient who was referred because of right scrotal mass. Using US, MRI, and CT, the mass was identified as a tumor. After local excision was requested by the patient, tumor was removed without rupture. Histopathology showed the tumor with mature adipose tissue missed with spindle cells and collagen fibers. Histopathology showed the tumor with mature adipose tissue along with spindle cells and collagen fibers. A hallmark of spindle cell lipomas is a positive stain of CD34, which was confirmed with this patient, along with vimentin and bcl-2 combined, confirmed the diagnoses of spindle cell lipoma (23).

Malignant neoplasms

Adenocarcinoma of the spermatic cord

Adenocarcinoma of the spermatic cord, though rare, presents with groin discomfort and swelling. The mass is palpable in the scrotum and the inguinal area (24). The most common occurrence of adenocarcinoma of the spermatic cord is due to metastasis from the stomach due to gastric cancer (24-26). Prognosis of patients with adenocarcinoma of the spermatic cord tends to be poor. However, in two cases presented by Kim et al., the follow-up was proven to be favorable (24). Histopathology reveals signet ring cells to identify the adenocarcinoma. US, MRI, and CT were all done to develop a proper image of the mass and determine if metastasis had prevailed. Multiple cycles of adjuvant chemotherapy was used with folinic acid, fluorouracil, and oxaliplatin (24).

Aggressive angiomyxoma of the spermatic cord

Minagawa et al. presented a case of a 37-year-old man who presented with a left lower abdominal mass. Clinical interpretation believed it to be an inguinal hernia. MRI revealed an inguinal mass extending from the left spermatic cord (27). In another case, CT was used to identify the mass (28). Surgical intervention was used to excise the mass and no recurrence was observed following the surgery (27). Aggressive angiomyxoma is more common in females than in males. It is extremely rare in males, more so in the inguinal region. It is locally aggressive but non-metastasizing (29). Patients commonly present with inguinal and scrotal swelling (27,30). Immunohistochemical results show emphasis on hormone receptors for estrogen and progesterone (31).

Lymphoma of the spermatic cord

Lymphomas of the spermatic cord are a rare occurrence (32). They can present as unilateral or bilateral masses with or without pain. These non-Hodgkin lymphomas are primary tumors that can be confused with testicular tumors that have extended into the spermatic cord (33). The most common form is diffuse large B-cell lymphomas reported in patients over 60, with a few reported cases of Burkitt’s lymphoma presenting in a 4-year-old boy and a 20-year-old young man (32-37). A recent study utilized different imaging modalities to characterize a primary spermatic cord tumor and distinguish it from other tumors presenting similarly; they reported preservation of normal funicular vessels, lack of fat within the lesion, as well as homogenous enhancement and intensity on MRI (33). Imaging is typically followed by radical orchiectomy and immunohistochemistry is utilized to confirm the diagnosis. Non-Hodgkin lymphomas stain
positive for CD20 among other markers and stain negative for CD3 and other markers (32-36). Generally spermatic cord lymphomas are known to have a poor prognosis but recent case studies show a good response to treatment (32). The patients were treated aggressively with multiple cycles of combination chemotherapy which includes rituximab, cyclophosphamide, doxorubicin, vincristine, and prednisone with some variation and additional therapy determined per case (32,34,35).

Malignant fibrous histiocytoma (MFH) of the spermatic cord

MFH is the most common soft tissue sarcoma in adults, however it is very rare for it to present as a primary spermatic cord tumor (38). It is generally a unilateral mass that is identified by imaging and requires a biopsy for a definitive diagnosis (39). Morphologically it is described to be a gray-yellow (sometimes white), elastic but hard mass (40-44). It is classified into four types based on microscopic examination: storiform-pleomorphic, giant cell, inflammatory, and myxoid of which the storiform-pleomorphic is the most common (42). The storiform-pleomorphic type is identified by abnormal spindle cells in a storiform or “star-like” pattern and giant cells with variation in size of nuclei (40-42,44). The inflammatory type shows the tumor cells with neutrophils permeating the tissue (43). Immunohistochemistry for MFH is positive for CD68 and vimentin and negative for cytokeratin and desmin with certain variations between cases (40-42,44,45). These tumors have a high chance of reoccurrence in the same area and/or metastasis (41,42,46). One case showed reoccurrence after 8 years with metastasis to the liver, lung, adrenal gland, bone, and mesenteric lymph nodes (46). The most common and recommended treatment option is surgical removal of the tumor using a wide local excision which can be followed up with radiotherapy (38,40,47). In some cases, chemotherapy agents such as ifosfamide and doxorubicin are used (41,43).

Malignant schwannoma of the spermatic cord

Malignant schwannoma or a neurilemmoma arises from the Schwann cells within the peripheral nerve sheaths (48,49). Though it is of unknown etiology, it is seen in patients with von Recklinghausen’s disease which can present as a neurofibromatosis type II gene deletion or alteration (48,49). This rarely occurs in the male genital area and there are very few cases reported of malignant schwannoma in the spermatic cord (50-52). Jiang et al. reports a case in a 20-year-old man who presented with a unilateral mass emerging from the spermatic cord which was hypoechoic on US (50). A biopsy is done for a definitive diagnosis; in this case, histology demonstrated spindle-shaped cells with elongated polymorphic nuclei (50). Immunohistochemistry for malignant schwannoma is positive for vimentin and S-100 protein and negative for smooth muscle actin—this is used to assist diagnosing the tumor (48,50). These patients typically have the tumor resected surgically; reoccurrence was not reported in the cases we reviewed (51,53).

Myxoid liposarcoma of the spermatic cord

Myxoid liposarcoma is the most common type of liposarcoma named for its myxoid matrix; it is also characterized to be a vascular tumor with low amounts of fat (54). It usually presents as an inguinal mass that is often mistaken for an inguinal hernia (54-56). Due to its unclear presentation, Abete et al. looked into imaging modalities which would assist in diagnosis (57). Their pre-operative diagnosis was based on a combination of MRI and US imaging; while the tumor had a cyst-like appearance on MRI, the vascularity of the tumor mass can be visualized on US (57). The tumor can also be diagnosed from its distinct histologically appearance post-biopsy, as discussed earlier. There is variability in the immunohistochemistry reported for these tumors (54,55). In most cases, the tumor was encapsulated with regular borders and therefore it was surgically removed without recurrence (55,57-59). There is one report of reemergence of the myxoid liposarcoma and metastasis to the liver after surgery (60).

Myofibroblastic tumor of the spermatic cord

Myofibroblastic tumors of the spermatic cord are identified by different names in literature: inflammatory myofibroblastic tumor (IMT), pseudosarcomatous myofibroblastic proliferation, and proliferative funiculitis (61-63). Case reports describe it to be a “painless scrotal mass” where US imaging can help detection but insufficient for an accurate diagnosis (61). There is controversy between a few earlier papers which claim IMT to be a benign mass while others refer to it as a low-grade malignant tumor (64-66). Fine needle aspiration or tumor biopsy is used to examine the tumor histology and immunohistochemistry (61,64). Histologically, the tumor is a composed of myofibroblasts and fibroblasts which appear spindle-shaped along with
permeating inflammatory cells (61,62,64,67,68). Coffin et al. described three different patterns which would present similar to a nodular fasciitis, a fibrous histiocytoama, or a desmoid (66). As a result, immunohistochemistry is very important in distinguishing the tumor from others on the differential diagnosis (61). While most tumors stain positive for vimentin and negative for S-100 protein, there is variation in markers such as SMA and ALK among cases (61). Coffin et al. stresses the importance of anaplastic lymphoma kinase (ALK) as a marker that assists in predicting the prognosis of the tumor—ALK is positively correlated with lower chances of metastasis (69). Patients were most frequently treated by wide surgical excision of the tumor and showed favorable results (61-64,67).

**Myxofibrosarcoma (MFS) of the spermatic cord**

MFS is also used for the myxoid type MFH (70,71). MFS in the spermatic cord is a soft tissue tumor that presents as a painful or painless scrotal mass (70,72). The tumor can be de novo or due to prior exposure to radiation (73). The tumor is composed of abnormal spindle cells within a myxoid background (70-73). Well-differentiated and dedifferentiated liposarcomas are often found to resemble MFS, however, they consist of lipoblasts which are absent in MFS, therefore immunohistochemistry can also be used to reach the correct diagnosis (74-76). MFS is similar to MFH as it is positive for vimentin and stains negative for S-100 and myogenin (71,73). Studies report high chances of recurrence as well as metastasis with MFS (71-73). These tumors are distinguished as high-grade or low-grade to determine prognosis and guide treatment recommendations (72,73). For all cases, surgical excision of the tumor is recommended. However, depending on the grade of the tumor, surgery will be preceded or followed by chemotherapy or radiotherapy (70,71,73).

**Osteosarcoma of the spermatic cord**

There are very few reported cases of primary extraskeletal osteosarcoma of the spermatic cord (77-79). Patients present with a painless mass that is generally calcified on a CT scan (77,78). The tumor is often described to have “bone trabecules” along with malignant cells, spindle cells, or giant cells which are multinucleated (77,78,80). In certain reported cases, this histologic appearance is combined with liposarcoma or other tumor types such as in cases of malignant mesenchymoma (80-82). There is no mention of immunohistochemistry of the tumor as it is distinguished histologically. The treatment for this tumor involves a wide surgical excision of the tumor along with tissues in nearby proximity; at times, this includes a radical orchietomy and ligating the spermatic cord to prevent recurrence (77,78,80). In certain cases, the surgery is followed by radiotherapy or chemotherapy (77). Beiswanger et al. reports a patient with the combined presentation of osteosarcoma of the spermatic cord and renal cell carcinoma; the patient showed no recurrence after surgical excision of the osteosarcoma and partial bilateral nephrectomies (78).

**Rhabdomyosarcoma of the spermatic cord**

Rhabdomyosarcoma is different from the other spermatic cord malignancies as it commonly presents at an earlier age, typically around age 4 and 18 (83-85). However, there are reports of rhabdomyosarcoma in the adult population (84,86,87). The tumor can arise from the normal muscle tissue in the male genital area or from mesoderm that has not differentiated (84,87,88). It is described to be a hard but painless mass found in the inguinal or scrotal region appearing grayish white and lobular when removed (84,85,88,89). Bouchikhi et al. discusses using imaging modalities to eliminate other possible tumors, however tumor histology is necessary for a definitive diagnosis (85). There are four histological types of rhabdomyosarcoma: embryonal, pleomorphic, alveolar, and mixed type (90). As the name suggests, histologically one would see rhabdomyoblast with multiple nuclei, mitotic activity, and other tumor cell characteristics (84,85). On immunochemistry, the tumor is identified by its positive staining for desmin and myogenin (84,85,88). These tumors are capable of metastasizing, therefore the abdomen and lungs are frequently examined upon diagnosis (85,86,88-90). The suggested treatment for this tumor in all cases has been a radical inguinal orchietomy followed by cycles of chemotherapy—typically VAC which includes vincristine, actinomycin, and cyclophosphamide (84-91).

**Summary**

Evidently, neoplasms are rare occurrences within the spermatic cord. Understanding the significance in rare findings are crucial factors in order for an accurate and timely diagnosis. Although the classifications of tumors appear to be painless and ignorable, a proper analysis may reveal possible linkages to other sources of irregularity such as discomfort or carcinoma. Therefore, emphasizing further
research on some of these overlooked results would lead to advancements in diagnostic and management strategies.

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Footnote

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