Paratesticular sarcomas: A report of seven cases

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Abstract. Primary tumors of the paratesticular region are rare, with paratesticular sarcomas constituting a major proportion of these tumors, particularly in the elderly. The paratesticular region consists of mesothelial, various epithelial and mesenchymal cells and may therefore give rise to a number of tumors with various behaviors. Defining the association between the paratesticular mass and the testicle, and differentiation between benign and malignant masses using radiology is challenging, therefore the mass is usually considered to be malignant and radical orchiectomy with high ligation is performed. The present study reports the cases of seven patients with tumors of the paratesticular region and presents the clinical and significant histological features of the tumors. In total, two patients suffered from dedifferentiated liposarcoma (DDLS), two exhibited leiomyosarcoma, two exhibited low-grade fibromyxoid sarcoma and one case of undifferentiated pleomorphic sarcoma was identified. Radical orchiectomy with high ligation was performed in five cases; simple orchiectomy was performed in the remaining case. A leiomyosarcomatous and epithelial membrane antigen (EMA) positive whorl pattern was observed during microscopy in the two DDLS cases. Additionally, one of the low-grade fibromyxoid sarcoma patients exhibited pleomorphism and mitosis in focal areas. To the best of our knowledge, only one instance of low-grade fibromyxoid sarcoma in a paratesticular location has been previously reported (7).

Introduction

As the paratesticular region contains various structures, including the epididymis, spermatic cord, tunica vaginalis and strong fat-ligament-muscle supporting tissues, it may give rise to a number tumor types with various behaviors (1). Tumor variability may also be due to the Wolffian duct origin of the testis appendages, including the spermatic cord. The most significant feature of the paratesticular region is that it is the origin of a small number of tumors with rich diversity.

The majority of the masses within the scrotum in adults are of testicular origin. Paratesticular masses account for 2-3% and sarcomas account for ~30% of all scrotal masses (1-4). The most common type of sarcoma is liposarcoma, followed by leiomyosarcoma (LMS), rhabdomyosarcoma (RMS), undifferentiated pleomorphic sarcoma and fibrosarcoma (1-6). To the best of our knowledge, only one instance of low-grade fibromyxoid sarcoma in a paratesticular location has been previously reported (7).

Determining the association between the paratesticular mass and the testicle, and differentiation between benign and malignant masses using radiology is challenging, therefore the lesions are usually considered to be malignant and radical orchiectomy with high ligation is used. The prognosis is often poor, as recurrence and metastasis are common, and the mechanism and outcome of regional lymph node resection, radiotherapy and chemotherapy is unclear. The present study reports seven cases of paratesticular sarcoma and emphasizes the significant clinical and histological features.

Case reports

Seven cases of paratesticular sarcoma diagnosed at the Pathology Department of the Istanbul Education and Research Hospital (Istanbul, Turkey) are retrospectively investigated. Hematoxylin and eosin and immunohistochemical staining of the cases were reevaluated and accurately diagnosed according to the recent World Health Organization classification (8). The...
clinical information of the patients was obtained from the patient files. Written informed consent was obtained from all patients.

All patients had been referred to the Urology clinic at Istanbul Education and Research Hospital with a growing scrotal mass. Excisional biopsy and simple orchiectomy were performed in cases three and four and the two patients were subsequently diagnosed with fibromyxoid sarcoma. Radical orchiectomy was performed in the other five cases. Cases one, two and three did not receive any additional treatment, whereas chemotherapy and radiotherapy was administered to case five. In cases four, six and seven, re-excision was performed due to recurrence, and chemotherapy and radiotherapy were administered following re-excision.

The clinical features are summarized in Table I. The macroscopic and histological characteristics of the cases were as follows: The lesions in cases one and two consisted of large yellow (lipomatous) and well-delineated areas, with occasional tan-gray colored (leiomyosarcomatous) areas. The diameter of the leiomyosarcomatous area was 8 cm in case one and 5 cm in case two. In addition, a few gray-colored nodules, the largest with a 1 cm diameter, were present in case two. A homologous pattern consisting of well-differentiated liposarcoma, comprising predominantly spindle cells, was present in each case (Fig. 1). The heterologous pattern consisted of a LMS and meningothelial-like whorl component in each case and a low-grade chondrosarcoma component in case one was also present (Figs. 2 and 3). Immunohistochemical staining of the cells revealed that the whorl pattern area was positive for epithelial membrane antigen (EMA; Fig. 4). A strong positive result for cluster of differentiation (CD)34 was observed in the spindle cells within the spindle cell liposarcoma areas, regarded as the homologous component (Fig. 5). The two cases were diagnosed as leiomyosarcomatosis and dedifferentiated liposarcoma (DDLS) containing whorl-pattern areas.

Cases three and four exhibited nodular, well-delineated tumors with surgical border invasion. Microscopy revealed the histology to be similar in the two cases, with the prominent features consisting of a collagen and myxoid zone, mixed with bland spindle-like fibroblastic cells and a whorl pattern, and arcades of curvilinear blood vessels (Figs. 6 and 7). The cellularity of the tumors varied from extremely low to moderate. Pleomorphism and mitosis were present in the nuclei in a focal area in case four (Fig. 8). Immunohistochemical staining for vimentin and MUC4 yielded a positive result in the two cases, together with focal CD34 and EMA positivity. The patients were diagnosed with low-grade fibomyxoid sarcoma.

The masses in cases five and six were well-circumscribed and nodular, with long bundles that were parallel or perpendicular to each other. The cell cytoplasm was strongly eosinophilic and the nuclei were generally spindle shaped, with one blunt end. Mitosis was not frequent, with 1-3 mitoses per 10 high power fields. Necrosis was observed in focal areas and pleomorphism was moderate. Immunohistochemical staining for smooth muscle actin and desmin yielded a positive result. The two cases were diagnosed as LMS.

Macroscopically, the mass in case seven had infiltrative borders, and, microscopically, was rich in spindle cells, with small bundles and storiform patterns. The tissue also contained large pleomorphic cells, with large eosinophilic cytoplasm in certain areas. Immunohistochemical staining yielded a positive result for vimentin only. This case was diagnosed as undifferentiated pleomorphic sarcoma.
Table I. Clinical findings and diagnosis of paratesticular sarcoma.

| Case | Age, years | Diagnosis                        | Tumor size, cm | Treatment                                    | Additional treatment | Follow-up time, months | Disease outcome |
|------|------------|----------------------------------|----------------|----------------------------------------------|----------------------|------------------------|------------------|
| 1    | 70         | Dedifferentiated liposarcoma     | 13.0           | Radical orchiectomy with high cord ligation | None                 | 14                     | Survival         |
| 2    | 38         | Dedifferentiated liposarcoma     | 13.0           | Radical orchiectomy with high cord ligation | None                 | 8                      | Survival         |
| 3    | 72         | Fibromyxoid sarcoma              | 7.0            | Excisional biopsy                            | None                 | 22                     | Mortality        |
| 4    | 63         | Fibromyxoid sarcoma              | 9.0            | Simple orchiectomy                           | Re-resection, CTh + RTh | 43                     | Mortality, recurrence with lung metastasis |
| 5    | 64         | Leiomyosarcoma                    | 6.5            | Radical orchiectomy with high cord ligation | CTh + RTh            | 21                     | Mortality, recurrence with lung metastasis |
| 6    | 68         | Leiomyosarcoma                    | 7.4            | Radical orchiectomy with high cord ligation | Re-resection, CTh + RTh | 18                     | Survival, recurrence |
| 7    | 46         | Undifferentiated pleomorphic sarcoma | 4.2            | Radical orchiectomy with high cord ligation | Re-resection, CTh + RTh | 44                     | Mortality, recurrence |

CTh, chemotherapy; RTh, radiotherapy.
Paratesticular sarcomas are rare and account for ~2% of all soft tissue sarcomas (9,10). No clear approach is available regarding their behavior and treatment due to their rarity. Liposarcoma and LMS are the most common sarcomas (1-6). RMS is more frequent in younger patients (6). The development of various paratesticular neoplasia is due to the differing complex structures in the region. In addition, embryological development of the spermatic cord, the most common tumor localization, and the other testicular adnexal structures from the Wolffian duct may be responsible for this histological diversity.

The histological features of the three cases diagnosed with LMS and undifferentiated pleomorphic sarcoma were typical and created no diagnostic problems. However, the two DDLS cases exhibited extremely rare histological features. Although the LMS and whorl patterns observed in the DDLS cases is not common, they have been described in previous studies (11-13). Leiomyosarcomatous areas were observed in each of the DDLS cases. The cases exhibited low-grade morphology and extremely low rates of mitosis, with mild pleomorphism in focal areas. The positive immunohistochemical staining for EMA in the whorl pattern areas was noteworthy and suggested perineural or meningeal differentiation. Meningothelial whorl-like morphology is less frequently observed than LMS. EMA was present in

**Discussion**

Paratesticular sarcomas are rare and account for ~2% of all soft
each case and, to the best of our knowledge, these are the first DDSLS cases positive for EMA to be reported.

A further feature of the DDSLS cases was the dominant spindle cell liposarcoma as a homologous component. The denomination of spindle cell liposarcoma remains controversial; certain authors use spindle-cell neoplasm or spindle cell lipoma-like neoplasm (14), and the recommended name for certain lesions was fibrosarcoma-like lipoid neoplasm, as determined in a report published by Deyrup et al (15). The authors who recommend these names do not classify these tumors as liposarcoma, however, lipoblasts were observed. The two DDSLS cases in the present study were rich in spindle cell lipoma-like (spindle cell liposarcoma) areas with strong cytoplasmic CD34-positivity, as well as rich in lipoblasts. These areas, which were the homologous component for DDSLS, should be classified as spindle cell liposarcoma.

Low-grade fibromyxoid sarcoma is a relatively rare sarcoma (16-17). The upper extremities and the torso are the most frequent locations, however, no paratesticular localization has been previously reported. Low-grade fibromyxoid sarcoma usually exhibits variable microscopic findings, with bland fibroblasts, whorls, linear sequencing and less cellular myxoid sections in certain areas (16). Mitosis and necrosis are rare. One of the present cases exhibited typical features of low-grade fibromyxoid sarcoma, while case four exhibited pleomorphism in focal areas that were also rich in mitoses.

The accepted treatment for paratesticular masses is radical inguinal orchietomy, including the surrounding soft tissues. No consensus with regard to regional lymph node excision has been reached, radiotherapy and chemotherapy. In the current study, radical orchietomy with high ligation was performed for the seven cases. Simple orchietomy and excisional biopsy were conducted for the two cases with fibromyxoid sarcoma. The two patients with fibromyxoid sarcoma succumbed to the disease within 22 and 43 months, respectively, as there was residual mass and the patients did not accept additional treatment. This emphasizes the importance of radical surgical treatments. No recurrence or metastasis was observed in the two cases of liposarcoma and the improved prognosis of liposarcoma compared with the other sarcomas, or the short clinical follow-up durations may have had an effect on this finding. As four of the seven cases succumbed to the disease and one remains alive with the sarcoma demonstrates the requirement for a multidisciplinary approach to the treatment of paratesticular sarcomas.

In conclusion, the paratesticular region consists of complex structures that can develop various neoplastic formations and patterns. Sarcomas comprise a significant part of paratesticular masses and may exhibit an aggressive clinical course. In older patients, paratesticular sarcomas must be considered for the differential diagnosis of scrotal masses, which do not exhibit a clear association with the testes. Furthermore, clinicians and patients must be informed about the high probability of local recurrence and distant metastasis in paratesticular sarcomas.

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