Giant paratesticular liposarcoma: A case report and review of the literature

ZUWEI LI1,2*, LIANG ZHOU1,3*, LIWEN ZHAO1, PEIJIE CHEN1,2, YIFENG LIU1, YU DING1, SHUOLEI SUN1, SHANGQI YANG1 and YONGQING LAI1

1Department of Urology, Peking University Shenzhen Hospital, Shenzhen 518036; 2Shantou University Medical College, Shantou, Guangdong 515041; 3Department of Urology, Guangzhou Medical University, Guangzhou, Guangdong 510006, P.R. China

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Abstract. Paratesticular liposarcoma is an infrequent tumor characterized by a growing, painless, inguinal or scrotal mass. Only about 200 cases have been reported as of yet in literature, however there are a few cases regarding giant paratesticular liposarcoma measuring over 10 cm. The disease may be commonly misdiagnosed prior to operation. Improper treatment tends to lead to local recurrence and distant metastasis. The current report presents a case of a 51-year-old patient with a large, painless right scrotum. Magnetic resonance imaging revealed a 7.8x5.8x10.4 cm nonhomogeneous space-occupying lesion of the right testis, which was firstly diagnosed as a spermatocytoma. Following this, a radical orchiectomy of the right testis was performed, however, it appeared to be a dedifferentiated liposarcoma, following histopathological examination and immunohistochemistry. Due to the large size of the tumor, it is significant to report the characteristics, diagnosis and treatment of the similar cases. The current study additionally presents a supplementary review of previously published cases in literature and focuses on discussion regarding the clinical characteristics, diagnosis, histopathology and immunohistochemical features and treatment of this disease.

Case report

In July 2017, a 51-year-old man, with a complaint of swelling of the right scrotum for 2 months, was admitted to the Department of Urology of our hospital. He presented with a painless and slow-growing fixed mass in the right scrotum without conspicuous promoting or alleviating factors. There are no other signs or symptoms. A rigid mass in the right scrotum, about 8 cm in maximum diameter, was the only positive finding of physical examinations. There are no specific abnormalities in the laboratory and imaging examinations (hemogram, urinalysis, stool routine, ESR, β-human chorionic gonadotropin, a-fetoprotein, mycobacterium tuberculosis antibody Ig-G, liver and kidney function tests and chest X-ray). However, magnetic resonance imaging (MRI) demonstrated a 7.8x5.8x10.4 cm nonhomogeneous space-occupying lesion of the right testis, which was firstly diagnosed as a spermatocytoma. Following this, a radical orchiectomy of the right testis was performed, however, it appeared to be a dedifferentiated liposarcoma, following histopathological examination and immunohistochemistry. Due to the large size of the tumor, it is significant to report the characteristics, diagnosis and treatment of the similar cases. The study was supported by the Ethics Committee of Peking University Shenzhen Hospital (Shenzhen, China) and written informed consent was obtained from the patient for the publication of the case details.

Introduction

Liposarcoma, a malignant tumor derives from mesodermal tissues, represents ~20% of all sarcomas. Paratesticular liposarcoma (PLS) is a rare condition. To the best of our knowledge, about 200 cases of PLS have been reported to date (1). Giant PLS is more rare with only a few cases having been reported (2-6). Due to the rarity of the disease, there is no standardized guideline as regards its incidence, diagnosis, recurrence and treatment (7,8). In this study, we present a case of a giant dedifferentiated PLS of the right testis with magnetic resonance imaging (MRI) measuring 7.8x5.8x10.4 cm and focus on the discussion about the clinical characteristics, diagnosis and treatment of this disease. Due to the giant size of this PLS, we report this case for the characteristics, diagnosis and treatment of the similar cases. The study was supported by the Ethics Committee of Peking University Shenzhen Hospital (Shenzhen, China) and written informed consent was obtained from the patient for the publication of the case details.

Correspondence to: Professor Yongqing Lai, Department of Urology, Peking University Shenzhen Hospital, 1120 Lianhua Road, Shenzhen 518036, P.R. China
E-mail: yqlord@163.com

*Contributed equally

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At 5-month follow-up, there is no evidence of local recurrence or distant metastasis.

Discussion

Liposarcoma, soft-tissue malignancy derived embryologically from mesodermal tissue, was first reported by Lesauvage in 1845 (9). They usually exist in the lower extremities and retroperitoneum (10,11). There are four histological subtypes in liposarcoma, which include well differentiated, dedifferentiated, myxoid and pleomorphic (12). PLS are rare neoplasm which compose approximately 12% of all liposarcomas and they originate in spermatic cord mostly followed by testicular tunics and epididymis (13). When the diameter of testicular tumor reaches more than 10 cm, such size will be called ‘giant’ (5). As far as we know, 200 or so cases of PLS have been reported up to date (14), and giant PLS are more rare with only a few cases having been reported (2-6). The incidence of PLS has a regional difference with the highest incidence being in Japan (7). The tumor attacked adult patients aged 50 to 60 years more frequently (15), though it occurred in patients with a range of 16 to 90 years of age on the basis of the current literature (16,17). PLS mostly present as a painless, slow-growing inguinal or inguinocrural mass and sometimes combine with a sensation of heaviness (9,14,16) and the occurrence of wrong diagnosis like scrotal lipoma, inguinal hernia and epididymitis before surgical intervention attribute to this clinical presentation (12,15). Most PLS are primary, but some can be metastasis from liposarcoma at other sites, such as thigh or the fatty tissue surrounding the testicle (18,19). Because of the insufficient number of literature on patients with PLS, no reliable standardized diagnosis and treatment guidelines have been made (14).

Ultrasonography (US), Computerized Tomography (CT) and Magnetic Resonance Imaging (MRI) are documented in the diagnosis of PLS (20,21). On US examination, PLS are identified as solid, heterogeneous solid, hypoechoic lesions, sometimes accompanied by colliquation if there is necrosis. However, US cannot always distinguish PLS from lipomas if the tumor is small or it is a well-differentiated PLS with homogenous fatty pattern, which makes PLS similar to lipoma (10,14,22). Compared to subcutaneous fatty tissue, CT usually demonstrates the tumor area with lower density. It may be helpful to establish tumor location, tissue characteristics,
staging and follow-up (8,16,23). MRI, the golden standard in staging soft tissue tumors, not only provides clear information on the tumor foci but also characterizes and delineates the degree of local tumor extension (9,14,20).

Diagnosis of PLS mainly depends on histopathology, immunohistochemistry and cytological features. A Critical histopathological analysis of dedifferentiated liposarcomas revealed that CD34 was negative in 9/11 cases; negative rate of s-100 was 92% (23/25); MDM2 was diffusely positive in well-differentiated areas and focally in dedifferentiated areas in the tumor with homologous dedifferentiation; SMA was positive in 2/8 tumors (24). Andrei et al proved that MDM2 and CDK4 were significative markers for confirming the diagnosis of well-differentiated liposarcoma (23). Histologically, differentiated sarcoma can be subdivided into five main subtype: Resembling pleomorphic malignant fibrous histiocytoma, fibrosarcoma, rhabdomyoblasosarcoma, myxofibrosarcoma and hemangiopericytoma (25,26). A total of 76% of dedifferentiated liposarcomas was high-grade (24).

Multimodality therapy was suggested by many researchers (27-29). There is a general consensus that radical orchiectomy with wide local excision and high ligation of the spermatic cord are the current standard treatment strategies due to frequent recurrence that associated with incomplete excision (7-9,20). Because the clinical presentation of PLS is similar to scrotal lipoma or groin hernia, Immediate radical procedure should be performed to avoid the high risk of local recurrence and involvement of worsening prognosis, when a suspicious PLS is diagnosed. It is important to prohibit spillage of malignant cells and acquire a more safe edge during the operation. A clinical research showed that the 3-year local-recurrence-free survival was 100% for negative margins compared with 29% for positive margins (30). Retropitoneal lymph node dissection is not recommended except for metastasis (7). It has been reported that occult local lesions were found at least a third of patients after operation (1), thus not only in dedifferentiated PLS, considered with a high rate of recurrence and metastasis, but also in other subtypes of PLS, adjuvant radiation is quite needed (9,14,31). Cerda et al (32) reported that five patients with spermatic cord sarcoma given adjuvant radiotherapy with a total dose of 54 Gy/27 or 30 fractions were found no recurrence in median 18 months of follow-up (range 6-28 months). However, whether radiotherapy should be used as postoperative routine therapy remains to be discussed because recurrent tumor after radiotherapy may be more aggressive (10). Some suggested that radiotherapy should be used for local control (8,10,14,30). There are no large studies with respect to the results of chemotherapy. A meta-analysis of 14 randomized clinical trials discovered that the improvement of recurrence and recurrence-free survival were attributed to chemotherapy (14,33). Some studies reported that we should attach importance to chemotherapy for high grade LPS (1).

Research report on prognosis of PLS is quite limited until now. A recent study local-recurrence-free survival was 76% at 3 years and 67% at 5 years (30). Another study about PLS revealed the 5-year survival rate was 75% and recurrence rate was 50-70% of all cases (14). Prognosis and overall survival rate vary in accordance with some risk factors, which include tumor grade, size, depth of invasion and histopathological classification (most important). The dedifferentiated types have a worse prognosis, but local recurrence rate will be smaller (14,34).

In conclusion, PLS represent a rarity of the tumor, characterized with slow growth, which are often misdiagnosed preoperatively. US, CT and MRI can redound to diagnose and differential diagnose, but the final diagnosis of PLS depends on histopathology and immunohistochemistry. When diagnosed or highly suspected preoperatively, radical orchiectomy with wide local excision and high ligation is the best treatment strategy and multimodality therapy is suggested. Long-term follow-up is recommended due to the risk of local recurrence and distant metastasis.

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