CASE REPORT

PRIMARY BREAST LEIOMYOSARCOMA IN MALE
A CASE REPORT

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ABSTRACT
Primary Sarcomas of the breast account for less than 1% of breast tumors. Leiomyosarcomas are less common. Only 51 cases have been reported in the literature, which only 4 cases in men. We present a new case of male breast leiomyosarcoma and discuss management and therapy of these unusual neoplasms. The confirmation of diagnosis was obtained by immunohistochemical study of surgical biopsy. The patient underwent a radical mastectomy and axillary lymphadenectomy. Adjuvant radiotherapy was indicated. Two years after surgery the patient is still in life without recurrence.

KEY WORDS: Breast – Sarcoma- Stromal tumors- Leimyosarcoma.

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INTRODUCTION
Primary sarcomas of the breast account for less than 1% of all primary breast tumours. [1, 2] The subtype Leiomyosarcoma (LMS) is less common. [3] Its current prevalence is 0.0006%. [4] This tumor arise from the mesenchymal tissue of breast and occurs usually in postmenopausal women. [5]. It occurrence in man is even rarer. We present a new case of male breast leiomyosarcoma and discuss management and therapy of these unusual neoplasms.

CASE REPORT
A 68-year-old man, without history of malignancy or prior radiation therapy on chest wall, was admitted for the management of painless tumefaction of the right breast growing gradually since 1 year, with inflammatory signs and invasion of adjacent skin. These symptoms were associated to anorexia and weight loss of 5 kilograms in 3 months. On Physical examination, we found a central ulcerated mass of the right breast (fig 1), attached to the deep plan of the chest wall with inflammatory signs and axillary lymphadenopathy. Computed tomography of the chest (fig 2) showed a tumor of 8x9 cm in diameter involving major pectoralis muscle without ribs involvement or pulmonary metastasis.

Surgical biopsy was performed, which histopathologic analysis revealed a proliferation of fusiform cells with atypical mitosis (9 to 10 by field) (fig 3). Immunohistochemical staining (fig 4 and 5) was positive intense and diffuse to Ki 67, SMA (smooth muscle actin), H’Caldesmon, negative to PS100 and to cytokeratin (AE1/AE3). A diagnosis of leiomyosarcoma was established. Initial workup including CT scan of whole body and bone scan was normal.

Figure 1: Ulcerated right breast mass
The patient underwent a radical mastectomy with resection of pectoralis muscle, axillary dissection, and immediate reconstruction with latissimus dorsi flap. Final histopathologic analysis of surgical specimen (fig 6) showed a necrotic tumor measuring 11x8x10 cm in diameter with free surgical margins (3 mm to the deep plan and 3cm to lateral margins) and confirmed the diagnosis of leiomyosarcoma grade II of FNCLCC.

DISCUSSION

Breast sarcomas are rare and represent less than 1% of breast tumors [1, 2]. Primary breast leiomyosarcoma are even rarer. In the scanned literature 51 cases have been reported prior to this observation [3, 5-10]. They were 47 women and only 4 men with median age of 56 years (24-83 years). There occurrence in men suggest the absence of endocrine factors in the genesis of these tumors [1, 11]. They develop from smooth muscles of areola or blood vessels [12]. Clinically we always found a painless mass growing progressively without invasion of deep structures of the chest wall [2]. In our case the major pectoralis muscle was massively invaded. Diagnosis is based on electron microscopy, histopathologic and immunohistochemical analysis of surgical biopsy performed before any treatment. These tumors usually express Smooth muscle actin, Desmin, and Vimentin and are negative staining for S100, cytokeratins, and epithelial markers [1, 2]. Mitotic activity of these tumors varies from 0 to 24 mitosis/10 high power field (HPF) with a mean of 11 mitosis/ HPF. [13] Surgery is the mainstay of treatment and the only guaranty of cure. The standard surgical procedure is radical mastectomy with surgical margins of at least 3 cm [12]. The risk of regional lymphatic extension is low and analysis of the case reports in the literature shows that lymphadenopathy usually corresponds to hyperplasia rather than a tumor, then axillary dissection is not mandatory [3, 5, 9, 12]. In our case axillary dissection has been done because physical exam found axillar macroscopic lymphadenopathy. These tumors likely spread by haematogenous way to lung, bone, liver, and central nervous system [14-16]. There is no consensus about adjuvant treatment by chemotherapy or radiotherapy, and the benefice of these additional treatments seems uncertain. [1-8] Radiation therapy ameliorated the local control in 3 cases previously reported in the literature. [17-19] In our case the deep surgical margin was insufficient, then postoperative radiotherapy was indicated to reduce risk of local recurrence. Pazopanib is a new treatment option for patients with metastatic non-adipocytic soft-tissue sarcoma after previous chemotherapy. The PALETTE study enrolled 369 patients, assigned to receive pazopanib (n=246) or placebo (n=123). Median progression-free survival was 4+6 months (95% CI 3+7-4+8) for pazopanib compared...
Primary breast leiomyosarcoma. [8]

CONCLUSION
Primary leiomyosarcoma of the breast is a rare tumour, even rarer in man. It can be a diagnostic challenge to clinicians as it has no specific clinical or radiological features. The final diagnosis is obtained by histopathological and immune-histochemical study. These tumors must be excised with a negative margin in order to obtain good results. The benefits of chemotherapy, radiotherapy, and hormonal therapy are still controversial. Radiotherapy may be indicated if the tumour size is large or if the resection is incomplete. Several studies need to be done to determine the prognostic factors.

AUTHORS’ CONTRIBUTIONS
The participation of each author corresponds to the criteria of authorship and contributorship emphasized in the Recommendations for the Conduct, Reporting, Editing, and Publication of Scholarly work in Medical Journals of the International Committee of Medical Journal Editors. Indeed, all the authors have actively participated in the redaction, the revision of the manuscript and provided approval for this final revised version.

REFERENCES