INTRODUCTION

Primary cutaneous leiomyosarcoma (PCL) is a rare malignancy. It accounts for 2 to 3% of all soft tissue sarcomas [1]. They can come from the erector smooth muscle of the hair at the origin of the dermal leiomyosarcoma also called cutaneous leiomyosarcoma or smooth muscles of the adipose tissue vessels giving the subcutaneous or hypodermic leiomyosarcoma.

We report a particular case of giant superficial leiomyosarcoma of the thigh with highlighting the diagnostic and management difficulties.

CASE REPORT

Figure 1a-b. Ulcerated tumor budding and infected on the inside of the thigh.
The result of the histological examination revealed a cutaneous sarcoma with spindle cell of high malignancy and the immuno-histochemical profile confirmed the diagnosis of a leiomyosarcoma. The extension assessment did not show secondary locations.

RESULTS

The patient underwent extensive excision with cutaneous margins of three to five cm and carried the Gracilis muscle deep down. The vascular pedicle and the large sciatic nerve were intact at a safe distance from the tumor (Fig. 3). After nearing the wound edges the loss of cutaneous substance is estimated at 20 * 15cm (Fig. 4). The exeresis limits were healthy on histological examination of the surgical specimen (Fig 5).

Figure 5. Macroscopic appearance of the operative specimen including the Gracilis muscle.

The patient is kept in the trauma department for local wound care; the monitoring is marked by good post-operative follow-up including no signs of infection. After budding of the residual wound bed, a thin skin graft was performed with very good aesthetic and functional results (Fig. 6). After one year of control we did not notice local recurrence.

Figure 6a. Wound cover by a thin skin graft. b. result after three months.
DISCUSSION

LCP occurs in people of all ages but with a peak between 60-70 years old. Superficial leiomyosarcomas have two subdivisions, the cutaneous and subcutaneous forms. Subcutaneous tumors have been associated with a higher risk of local recurrence and distant metastases compared to the cutaneous form [2].

The clinical aspect is not specific with a wide range of differential diagnoses: squamous cell carcinoma, achromic melanoma, and basal cell carcinoma [3]. LCP is generally considered a solitary nodule; the nodule may be lobulated, pedunculated or umbilicated with a surface that may be smooth, indurated, ulcerated, scaly, verrucous or hemorrhagic [4]. In our patient, we observed the solitary and nodular character with ulcerated smooth surface. It is a fast growing mass and the prognosis is poor if the mass size is greater than five cm [4]. For our case the size is voluminous, to our knowledge; no case in the literature has been reported with such a volume. The risk of relapse is not zero even in the case of R0 exeresis of dermal LCPs which is not very recurrent [5]. This is perhaps why some authors argue for a wide excision with margins of three to five cm and a depth that includes the subcutaneous tissue and fascia [6].

Adjuvant therapies include radiotherapy, chemotherapy [4]. However, LCP has been reported as radio-resistant; also chemotherapy with doxorubicin has failed in some cases [1].

CONCLUSION

Primary cutaneous leiomyosarcoma is a rare entity whose clinical presentation may seem nonspecific, making diagnosis difficult. Anatomo-clinical correlation and immunohistochemical examination are mandatory for the definitive diagnosis.

Surgical excision with wide margins of safety is the standard treatment. Other therapeutic methods, such as radiotherapy or chemotherapy, are described but without significant benefits. Despite radical surgical treatment, because of recurrence rates, the prognosis remains poor. We recommend long-term follow-up of patients to prevent recurrence.

REFERENCES