Giant Mixed Lipoma/Liposarcoma of the Thigh: Diagnostic Approaches and Surgical Management

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Abstract
A 58 years old male with a subcutaneous neoformation, localized at the medial surface of the left thigh, of about 30 cm in diameter, and which has grown larger, particularly during the last year is described. Clinical signs inclined towards a diagnosis of a soft-tissue neoplasm such as giant lipoma. Gigantic lipoma of the thigh are rare. These lesions are of interest because of their risk of recurrence following surgical excision, as well as their potential risk of malignant transformation. We present a case of giant, deforming lipoma with 5% of the mass resulted compatible with dedifferentiated grade III liposarcoma of the thigh, with emphasis on the diagnostic difficulties encountered, and the surgical management of the case.

Keywords: Giant lipoma; Mixed liposarcoma; Open surgery; Adipose tumors

Introduction
The most common and frequent subcutaneous tumour is well-differentiated encapsulated lipoma composed of mature adipocytes (mature lipoid cells). This neoplasm, with a prevalence rate of 2,1/1000 [1,2] is a slow growing, with a somewhat soft consistency. The ethiology of lipoma is still unknown. Lipoma can be casual, or they may be hereditary. Theoretically metabolic/endocrine or genetic disorders may be involved [3]. Traumatic events have been implicated in the development of lipomas [4]. Lipomas frequently are small, weighing just a few grams, and they are usually less than 2 cm in diameter. They are usually movable but sometimes fixed by overlying adjacent tissues, which they may displace. However, these lesions can weigh over 200g and measure over 10cm in diameter [5].

Study Aim
This study describes our case management of a giant liposarcoma of the thigh in July 2005.

Methods
Surgical treatment
The patient was operated under general anaesthesia in a dorsal-recumbent position.

The mass, as well as the overlying skin, and the biopsy tracts were removed. To reach the mass, the blood vessels were moved aside, the fascia was incised circumferentially, the mass was identified and isolated, the sciatic nerve was moved aside, and the mass removed intact, together with the gracilis muscle. The surgical resection consisted of a 35x25x27 cm soft tissue specimen. We left a macroscopically clean margin. Two lymph nodes were identified and removed from the fascia. To cover the femoral vasculo-nervous complex, the nearby muscles were converged and a large distal, pedicled, fascio-cutaneous flap was harvested from the thigh. The donor site was covered with a dermatome-obtained skin graft. A drainage tube was placed, in aspiration, under the flap. Intra-operative antibiotics were administered and continued post-operatively. Heparin, 4000 i.u. a day was administered. At the end of the operation, the haemoglobin value decreased 1.8 g/dl to 10.8 g/dl with a haematocrit value of 32.%. On the fifth post-operative day an infection of the apex of the flap (which was otherwise perfectly vital) arose. Microbiological culture study found a Pseudomonas Aeruginosa. Using an appropriate antibiotic therapy, in association with frequent dressing changes, the infection was successfully treated. Only a slight wound distal dehiscence of the flap healed by second intention.

Figure 1a and b: Giant lipoma-liposarcoma of the left thigh in a 58 y.o caucasian male: preoperative view.

Figure 2: Magnetic resonance (MR) of the left thigh showed a voluminous, multi-lobed mass (27 X 20 X 22 cm), with distinct and regular margins, non-infiltrating, but clearly dislocating the adjacent muscle layers.
of the neurovascular complex was also dislocated laterally and forward. These results suggested a diagnosis of liposarcoma.

However, a preliminary biopsy, performed in a random area of the mass, resulted as lipoma. Subsequently, multiple MRI guided biopsies were performed which resulted as “compatible with well-differentiated liposarcoma (atypical lipoma)”. Further testing such as Chest CT scan and abdominal sonogram excluded distant metastases. On the advice of Oncology and Radiotherapy specialists, a wide surgical excision to permit a complete histological examination of the neoplasm and the study of its biological characteristics was performed by our team, together with an orthopaedic team.

Results

Histologic findings

The definitive histologic examination evidenced “dedifferentiated grade III liposarcoma”, according to the FNCLCC system. The margins were clear and the two lymph nodes showed reactive hyperplasia.

The specimen weighed 8 kg and only 5% of the mass was found to be liposarcoma (Figure 3).

The adipocytes showed focal nuclear atypia and hyperchromasia in the presence of stromal cells, multinucleated, especially in fibrous septa, numerous lipoblasts mono and multinucleated. Immunohistochemical studies showed positive expression for vimentin, S-100, CD30, CD45, desmin, myogenin, smooth muscle actin. Atypical lipomatous tumor subtype liposarcoma well-differentiated adipocyte (lipoma like) (Figure 4).

Patient management

Post- operative total body CT scan was negative.

The patient was treated with a cycle of external radiotherapy, following surgical excision. At 96 months follow-up after surgery no local or distant metastases were found (thigh MRI plus TB CT scan). The patient reports a good quality of life and satisfaction with the outcome (no residual discomfort, no neurologic defect or recurrence or adenopathy), (Figure 5a and 5b).

Discussion

According to Sanchez et al, in order for a lipoma to be called “giant”, the lesion must be at least 10 cm in one dimension or weigh a minimum of 1000 gr. [5]. One of the largest cutaneous lipomas,
reported by Brandler [6] in 1894, was 22.7 Kg, and located on the left scapular region, in a male, 26 years-old patient. Lipomas can grow in many different tissues or organs [1], and they have been found in the extremities, the gastrointestinal tract, the mediastinum, the heart, the brain, and spinal cord.

The malignant transformation of lipoma to liposarcoma is relatively uncommon. A few reports suggest that large tumours (> 10 cm in size) are more at risk of containing sarcoma cells [2,3], especially in the presence of sudden rapid growth [7]. Liposarcoma is the most common malignant mesenchymal neoplasm in adults, classified in 5 subtypes, including well-differentiated, dedifferentiated, myxoid, pleomorphic and mixed liposarcoma. There are also reports that the intramuscular location of a lipoma is a risk factor for malignancy [2,3]. The lower extremities are one of the most common sites for liposarcomas [8,9]. In this site, when diagnosis of liposarcoma is sure, Campbell et al recommends wide excision or amputation if needed [10]. They have reported a high probability of recurrence (91%) secondary to conservative initial treatment. Some authors have reported that postoperative adjuvant radiotherapy lengthens the median survival rate when surgical excision has not been adequate [7,8,10].

Giant lipoma treatment can be by open surgical excision or by liposuction, but it is very important to choose the most appropriate operative technique, considering the possibility of malignant transformation [12].

Fine needle aspiration biopsy and core biopsy usually do not provide sufficient material to allow the pathologist to differentiate between lipoma and liposarcoma [11]. Incisional biopsy is superior especially when performed in selected areas using MR imaging [13,14].

In our case, the first incisional biopsy wasn't enough to allow correct diagnosis. We suspected a malignant transformation considering the MR images, the size and fast grow. A second incisional biopsy guided by the MR images, allowed the diagnosis of well differentiated liposarcoma. We then excluded distant metastases using CT scan and abdominal sonogram. We then consulted oncological and radiotherapy specialists who suggested a wide surgical excision with a complete histological examination of the whole neoplasm.

According to the literature we chose open radical surgery to removal the whole, capsulated neoplasm, the overlying skin of the external aspect, and the gracilis muscle at the inner aspect.

Currently, the patient is disease free at 8 years follow-up, without surgery associated morbidity and with an optimal satisfaction of the surgical outcome.

**Conclusions**

In our opinion open and wide surgical excision, respecting oncological criteria of radicality, is the mandatory method of giant lipoma treatment. Surgical excision allows a complete removal of the macroscopically visible neoplasm, permitting a complete histological evaluation of the tumour to exclude tissue malignant transformation. Giant lipomas present an high risk of coexisting internal liposarcoma areas, these areas are very difficult to be identified with preoperative studies.

**Competing Interests**

The authors declared that they have no competing interests.

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