Desmoid Tumor of the Thigh with Multiple Recurrences
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Abstract

**Background:** Desmoid tumors are rare neoplasms of uncertain etiology arising from fascial or deep musculo-aponeurotic structures. Although with benign histological appearance and no metastatic potential, desmoid tumors are locally aggressive tumors with a high rate of local recurrence.

**Case Report:** The case of a 47 years old woman repeatedly operated for a recurrent desmoid tumor of the right thigh is presented. The initial radical excision was followed by adjuvant radiotherapy but local recurrence was diagnosed one year after. The pathology report revealed aspects of desmoid tumor. The patient was reoperated and subsequently referred for chemo- and hormonal therapy. A second recurrence occurred 20 months later and the patient was again operated with microscopic tumor free margins and positive long term outcome.

**Conclusion:** Desmoid tumors are benign tumors with unpredictable natural history. Best management involves a multidisciplinary approach. Concerning diagnosis, the best approach is performing a MRI examination. Wide surgical resection with adjuvant radiation therapy remains the main treatment option for local control.

Keywords: Desmoid Tumor; Surgery; Recurrence

Introduction

Desmoid tumors are rare neoplasms of uncertain etiology arising from fascial or deep musculo-aponeurotic structures that may occur at any age, but usually in young adults with peak prevalence between 25 and 35 years [1-3]. In adults, the tumor has a predilection in premenopausal women. Incriminated etiological factors are trauma or local surgery history, genetic factors like inherited mutation in APC gene (adenomatous polyposis coli) and high estrogenic states, including pregnancy [4]. Characterized by a benign histological appearance and no metastatic potential, desmoid tumors are locally aggressive tumors with a high rate of recurrence [5]. They may be localized in the abdominal wall, the bowel, and the mesentery (associated with familial adenomatous polyposis) or in extra-abdominal sites, such as the trunk and the extremities. The incidence of desmoid tumors ranges from 2 to 4 per million and almost half of them occur in the extremities and trunk [6].

Case Report

A 47 years old woman was hospitalized in our clinic for a swelling of the upper third of the right thigh, gradually increasing in size in the last 3 months. She had a history of an incomplete tumor excision on the same region, performed in another surgical unit 1 year ago. The pathological report showed at that time a fibroma with an important adipose structure. The local physical examination showed a tumoral mass located on the external face of the right thigh, under an old 10 centimeters long scar. The tumor was 8×4 cm, of firm consistency, painless, relatively fixed and seemed to infiltrate the subcutaneous tissue of the described area (Figure 1).

![Figure 1: Preoperative aspect.](image)

No motor, sensitivity or vascular alterations of the right lower limb were found. The CT scan revealed an expansive irregularly outlined tumoral mass on the upper third of the right thigh and gluteal region topography. The superior extremity of the tumor reached 1 centimeter above the upper bound of the right acetabulum and infiltrated the large and middle gluteal muscles. The tumor also invaded the semitendinosus, semimembranosus and right biceps femoris muscles without bone lesions. Under general anesthesia a wide tumor excision was performed with macroscopic tumor free margins (Figure 2).
The postoperative follow-up showed no signs of recurrence after 6 months. As recommended by the oncology board, adjuvant therapy with tamoxifen. After 20 months the patient was hospitalized for chemotherapy with metothrexate and vinblastine and antihormonal therapy with sulindac. A long-acting analog of indomethacin were associated with the partial and complete responses in several nonrandomized retrospective studies, either alone or in combination with hormonal agents such as tamoxifen. However, local control remains difficult. Radiation therapy with doses of 50–60 Gy is a viable alternative to surgery and a useful adjunct to incomplete resection of primary extra-abdominal desmoid tumors for the control of residual disease [8-11]. Radiation therapy alone can be used where surgery might lead to major morbidity and loss of function as well as for patients who have an inoperable tumor or gross residual disease after operative debulking [9].

The predisposition of aggressive fibromatosis to locally recur makes it difficult for the surgeon to grossly identify the true extent of the tumor extent and the progression of the disease before and after treatment. It may also be helpful in differentiating tumour progression from post-surgical fibrosis [7]. Because multicentric and recurrent lesions tend to occur within the same limb or anatomic region, the MRI scanning of the entire extremity will be done once the diagnosis is made.

As far as the therapeutic options are concerned, surgery is the treatment of choice for extra-abdominal desmoid tumors. Wide excision with free tumor marginal resection is the goal standard of primary treatment. Re-excision for treating the recurrent disease is preferred by most authors, resulting in a cure rate similar to that of the primary surgical resection [8]. However, local control remains difficult. Radiation therapy with doses of 50–60 Gy is a viable alternative to surgery and a useful adjunct to incomplete resection of primary extra-abdominal desmoid tumors for the control of residual disease [8-11]. Radiation therapy alone can be used where surgery might lead to major morbidity and loss of function as well as for patients who have an inoperable tumor or gross residual disease after operative debulking [9]. The use of antihormonal therapy for the treatment of desmoid tumors is based on epidemiological observations for example, higher incidences of desmoids during and after pregnancy and reports of spontaneous tumor regression after menopause [12]. Studies have shown that virtually all desmoid tumors express nuclear estrogen receptor-β, but only a small subset of patients respond to antihormonal therapies [13]. Because COX-2 seems to play a role in the pathogenesis of desmoid tumors, treatment with Non-Steroidal Anti-Inflammatory Drugs (NSAID) that inhibits COX may be effective [14]. A variety of other NSAIDs such as indomethacin and sulindac, a long-acting analog of indomethacin were associated with the partial and complete responses in several nonrandomized retrospective studies, either alone or in combination with hormonal agents such as tamoxifen. In contrast, in cases of an unresectable, rapidly growing and/or symptomatic and/or life-threatening desmoid tumor, traditional cytotoxic chemotherapy mainly with metothrexate and vinblastine may be the treatment of choice. Loco regional chemotherapy in the form of isolated limb perfusion for patients with locally advanced tumors is another alternative to systemic chemotherapy in patients with limb desmoids. Melphalan and recombinant human tumor necrosis factor-α are used as therapeutic agents with overall response rates of up to 80% [16]. The recurrence rates after wide local excision is reported as more than 40%, related to section margins, extra-abdominal localization of the tumor and age [8,17,18]. The predisposition of aggressive fibromatosis to locally recur is related to its infiltrative nature, the lack of pseudo capsule and possibility of diffusion along muscle fibres and fascial planes which makes it difficult for the surgeon to grossly identify the true extent of disease [19]. This may justify the high recurrence rate of the disease after adequate surgery and may also explain the distant recurrence on the same hip occurred to our patient.

The patient was reoperated and a radical surgical excision with microscopically free margins (frozen section examination) was done. The postoperative follow-up showed no signs of recurrence after 6 months.
examination. Wide surgical resection with adjuvant radiation therapy remains the main treatment option for local control.

Conflict of Interests

Authors have no conflict of interests to disclose.

References