Bilateral Breast Fibromatosis in a 41 Years Old Woman

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Authors’ contributions

This work was carried out in collaboration between all authors. Author AF evaluated the case and wrote the first draft manuscript with author JPE. Authors GEO, JMMN and CK contributed in data collection and literature search. Author JLEO contributed in manuscript preparation, editing and review. All authors read and approved the final manuscript.

ABSTRACT

Breast fibromatosis is a rare benign tumor of soft tissues with local spread, frequently recurrent but with no metastatic evolutions. We present a case diagnosed in a 41 years old patient referred to a Douala private medical center in February 2012, presenting bilateral breast tumor. Histological analysis of the excised samples showed fibroblastic proliferation without atypia and metastasis which reappeared 15 months after the initial intervention in the left breast. A mastectomy and lymph nodes curettage has shown resolution of the symptoms.

Keywords: Fibromatosis; histopathology; breast; Cameroon.

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1. INTRODUCTION

Fibromatosis is a rare tumor of the breast in females between puberty and menopause. This tumor can arise from the fascia musculoaponeurotic of the chest wall or from fibroblasts and myofibroblasts of the mammary parenchyma [1,2]. The neoplasm is characterized by being locally aggressive and frequent recurrence without metastasis [3,4]. The mammary fibromatosis represents only a particular aspect of these tumors; its frequency is less than 0.2% of the primitive tumors of the breast and 4% of the extra-abdominal fibromatosis [3]. Bilateral desmoid tumors have been reported in about 4% of patients [3]. The clinical presentation is a palpable mass that is sometimes associated with skin modification [5]. The diagnosis is based on histological examination of the biopsy or surgical specimen [1,2,6]. We are reporting a case of bilateral desmoid tumor of the breast. The informed consent from the patient for this study was obtained.

2. CASE PRESENTATION

A 41 years old female patient was referred to a private medical center in Douala town because of bilateral breast masses, a mass in the inferior external quadrant on the right and mass in the upper external left breast. In the two cases, the masses were mobile, painless and they appeared polylobular with no apparent changes in skin pigmentation and without any suspecting lesion.

Mammography showed large opaque masses with regular contours in both cases. The right breast measurement was 10x6x4 cm and the left was 11x7x5 cm. The cytological examination of the tissue aspirate taken under ultra-sound guidance showed abnormal cells.

The excised samples measured a 9x7x5 cm and weighted 58 g in the right breast. In the left, we obtained the sample measurement of 7x4x3 cm and 46 g in weight (Fig. 1). There masses were of firm consistency, well circumscribed with smooth and whitish surface homogenous on different sections. The specimens were treated according to the technical routine with fixation in 10% formalin, staining in eosin-haematein and interpretation under optical microscope.

In histological examination, the lesion was characterized by well and differentiated fibroblastic proliferation, which was rich in collagen without atypia, no images of mitosis. In some areas, simple mesenchymatous tissue and few blood vessels were observed (Fig. 2).

3. DISCUSSION

Desmoid tumors (DT) also known as aggressive fibromatosis (AF) constitute of a rare fibroblastic proliferative disease. They may occur in any musculoaponeurotic or fascial tissue [7]. These tumors generally are divided by anatomic designation as extra-abdominal, abdominal, or intra-abdominal and the most common locations are around the limb girdles or the proximal extremities, the abdominal wall and intra-abdominal or mesenteric [8,9]. The mammary fibromatosis is a rare lesion; bilateral type have been reported in up to 4% of patients and mostly it occurs in the breasts of women between 13 to 80 years with an average of 43 years, regardless of their ethnicity [2,3,5]. In our presented case, the breast location, the bilateral position of the tumor and the age of the patient, confirm in the data of the literature makes it a rare and interesting case.
Clinically, our patient presented bilateral breast masses, a mass in the inferior external quadrant on the right and another mass in the upper external left breast, associated with painless and no apparent changes in skin pigmentation with no suspecting lesion. Wongmaneerung et al. described two cases of bilateral breast fibromatosis with pain (of the patients) and nipple retraction, the presence of skin involvement and invasion by the tumor (in another patient) [5]. Mehdi et al reported a case with ulceration of all the nipple-areolar plate [1]. That means the clinical manifestations of breast fibromatosis are variable, making each case exceptional.

Although etiopathology is not fully understood, some theories had been proposed. Most desmoids arise sporadically, slightly more in women than in men, with some DTs related to pregnancy and trauma, and others associated with hereditary cancer syndromes [10]. Desmoid tumors are results of deregulation of connective tissue growth. Increased nuclear expression of β-catenin, a protein responsible for regulation of gene expression, proliferation and survival, is the characteristic feature in sporadic DT [10].

Antecedent trauma, often surgical, has been noted at the site of the DT in approximately 25% of cases [11]. Estrogen receptors (ER) were observed in 33% of all DT examined, with an equal incidence in males and females and with antiestrogen binding sites found in 79% of samples, including some which were ER negative [11]. Implant-associated breast desmoid tumors may also occur [12].

In this case, mammography showed large opaque masses with regular contours in both breast. In the literature, Mammography show typically high density, speculated and stellate tumor without microcalcifications, often indistinguishable from carcinoma [4,13]. The lesion sometimes appears as lobulated mass or mass with no outlines, but can still be well defined. In some rare cases, the lesion has no mammographic presence [14,15]. Although Ultrasound appearances are various, they are not specific. Demoid tumours could be poorly marginated; it can be irregular shaped hypoechoic mass or small-sized lesions that are homogeneous with regular outlines [15,16]. By some authors, the magnetic resonance imaging
(MRI) and the scanner are important for the estimation of a possible parietal invasion; MRI can also play an invaluable role in preoperative diagnosis and planning. Desmoid tumors are typically isointense on T1-weighted imaging and demonstrate low to high signal intensity on T2-weighted imaging. After the injection of chelated forms of gadolinium, in T1-weighting, and fat signal saturation, a heterogeneous raise in the ill-defined outlines should be observed [12,15].

Based on available reports in the literature, fine needle aspiration cytology of the entire specific may reveal some important information in patients with breast fibromatosis [17]. A cytopnction could confirm the diagnosis by isolating fibroblasts with minimal atypia; however, it is mostly a little contribution [2]. In our case, the cytological examination only showed abnormal cells.

The histological examination remains the basis for the diagnosis of this pathology. For our patient, histologically they found well and differentiated fibroblastic proliferation, which was rich in collagen without atypia, with no images of mitosis. In some areas, simple mesenchymatous tissue and few blood vessels were observed. According to the literature, the tumor usually has a poorly circumscribed pattern and is composed of proliferating stellate to spindle cells arranged in long fascicles or whorling patterns with bland nuclear features and dense keloid-like collagen in areas [10]. Ultrastructural studies indicate that the spindle cells have features of both fibroblasts and myofibroblasts. Architecturally, the tumor cells are typically arranged in long, sweeping fascicles, and vague whorls in a background of eosinophilic, collagenized stroma with prominent thin-walled vessels. The stroma can also show myxoid features, reportedly more common in the breast and the mesentery and can also be seen in other more common sites [11].

Immunohistochemistry, specifically β-catenin, and more recently, molecular diagnostics can play an important role in its diagnosis. The presence of actin and vimentin is useful for the diagnosis of desmoid tumor. Desmin is rarely positive, S100 and CD34 are usually negative; β-catenin nuclear staining is an option for diagnosis, which may be only focally positive. Sporadic DTs are commonly associated with somatic mutations of the codons 41, 45 of exon 3 of the beta-catenin gene (CTNNB1). As such, antibody of β-catenin is useful in distinguishing desmoids from its histologic mimics, which generally lack this feature [5,10,11,18].

The differential diagnosis of desmoid tumor vary from benign reactive lesions such as a hyperproliferative scar, to a more sinister fibrosarcoma [11].

The treatment of this patient was surgical. Radiotherapy was not directly demanded. The management of breast fibromatosis includes surgical excision with clear margins, Systemic therapy and radiation therapy [6,11]. Surgical therapy remains the cornerstone of desmoid tumor management. The mastectomy can be indicated for vast fibromatosis or for too large recurrences. In case of parietal invasion, the surgery can be extremely decaying going as far as taking away the breast, the pectoral muscles, the thoracic wall, and the parietal pleura [15]. Obtaining microscopically negative margins (R0 resection) is the preferred objective in oncologic surgery, including resection of desmoid tumors. Not all positive margins were recurrent; the true impact of surgical margin negativity as well as the role of adjuvant therapies i.e., radiation, antiinflammatory drugs, antiestrogen agents, and cytotoxic chemotherapy in preventing tumor recurrence remain uncertain [11]. Recurrent disease has been observed in negative margins, some authors proposed the algorithm of a more conservative wait-and-see approach in a less critical site with an asymptomatic lesion [19]. If the lesion progresses by RECIST (response evaluation criteria in solid tumor) criteria, then treatment is needed. Desmoid tumors respond to systemic medical therapies, from antiinflammatory drugs to standard chemotherapy agents. Liposomal doxorubicin, doxorubicin-based regimens, methotrexate in combination with vincristine and vinorelbine have demonstrated notable responses in this disease. Molecular-targeted agents, antiinflammatory drugs, hormonal agents, and interferons have all shown various degrees of activity in DT. However, the mechanisms through which these compounds conduct their antitumor activity are not completely understood [20].

Radiotherapy can be used in patients who have recurred after initial surgery and as primary treatment in those patients who are medically inoperable or those to whom resection presents unacceptable morbidity [21]. Some studies have shown a beneficial effect of the radiotherapy with 60% of decreasing cases non-accessible to the surgery. Therefore, the radiotherapy could be curative in recurrence cases when the surgery is impracticable [22,23]. The use of radiotherapy must be considered carefully in light of the
potential long-term side effects, including the risk of secondary malignancy, particularly in younger patients. Treatment decisions are individualized to each patient depending on a range of patient and tumor characteristics [24].

4. CONCLUSION

Breast fibromatosis is a benign affection which can be mistaken clinically, radiologically and cytologically as a malignant lesion. The bilateral form is extremely rare. A well-coordinated, multidisciplinary approach involving the input of surgical and nonsurgical specialists is needed to develop an individualized treatment strategy appropriate for each specific patient.

ETHICAL APPROVAL

It is not applicable.

COMPETING INTERESTS

Authors have declared that no competing interests exist.

REFERENCES

1. Mehdi E, Jaouad K, Mouna A, Abdellah B, et al. Primary fibromatosis of the breast in a 13 years old girl. Open Journal of Obstetrics and Gynecology. 2012;2:206-209.
2. Croce S, Letourneux C, Dale G. The mammary fibromatosis: A benign lesion relatively unknown. Gynécologie-Obstétric & Fertility. 2009;37:442-446.
3. Taylor TV, Sosa J. Bilateral breast fibromatosis: Case report and review of the literature. J Surg Edu. 2011;68(4):320-325.
4. Neuman HB, Brogi E, Ebrahism A, Brennan MF, Van Zee KJ. Desmoid tumors (fibromatoses) of the breast: A 25-year experience. Ann Surg Oncol. 2008;15(1):274-280.
5. Wongmaneerung P, Somwangprasert A, Watcharachan K, Ditsatham C. Bilateral desmoid tumor of the breast: Case series and literature review. International Medical Case Reports Journal. 2016;9:247-251.
6. Bonvalot S, Eldweny H, Haddad V, Missenard G, et al. Fibromatoses extra-abdominales primitives: Certaines patients peuvent-ils éviter la chirurgie ? e-mémoires de l'Académie Nationale de Chirurgie. 2007;6(2):72-80.
7. Goldblum J, Fletcher JA. Desmoid-type fibromatoses. In: Fletcher CDM, Unni KK, Mertens F (Eds) World health organization classification of tumours. Pathology and Genetics of Tumours of Soft Tissue and Bone Lyon: IARC Press; 2002.
8. Easter DW, Halasz NA. Recent trends in the management of desmoid tumors. Summary of 19 cases and review of the literature. Ann Surg. 1989;210(6):765-769.
9. Brown CS, Jeffrey B, Korentager R, Hughes K. Desmoid tumors of the bilateral breasts in a patient without Gardner syndrome: A case report and review of literature. Ann Plast Surg. 2012;69(2):220-222.
10. Escobar C, Munker R, Thomas OJ, Li DB, Burton VG. Update on desmoid tumors. Annals of Oncology. 2012;23:562-569.
11. Litchman C, editor. Desmoid tumors. New York (NY): Springer; 2011.
12. Jeong WS, Oh TS, Sim HB, Eom JS. Desmoid tumor following augmentation mammoplasty with silicone implants. Arch Plast Surg. 2013;40(4):470-2.
13. Lee SM, Lee JY, Lee BH, Kim SY, Joo M, Kim JI. Fibromatosis of the breast mimicking an abscess: Case report of unusual sonographic features. Clin Imaging. 2015;39(4):685-688.
14. El Demellawy D, Herath C, Khalil M, et al. Fibromatosis of the male breast with concurrent florid- type gynecomastia: Report of a case and review of literature. Pathology—Research and Practice. 2011;207:306-309.
15. Mesurolle F, Ariche-Cohen M, Mignon F, et al. Unusual mammographic and ultrasonographic findings in fibromatosis of the breast. European Radiology. 2011;11:2241-2243.
16. Rammohan A, Wood JJ. Desmoid tumour of the breast as a manifestation of Gardner's syndrome. International Journal of Surgery Case Reports. 2012;3:139-142.
17. Zanella M, Falconieri G, Della D. The value of fine needle aspiration cytology in breast fibromatosis. Study of two new cases and review of the literature. The Breast Journal. 1999;5(4):264-268.
18. Aitken SJ, Presneau N, Kalimuthu S, et al. Next-generation sequencing is highly sensitive for the detection of beta-catenin mutations in desmoid-type fibromatoses. Virchows Archi. 2015;467(2):203-210.
19. Al-Yusuf R, Fakhro AR, Alkhaznah A. Breast fibromatosis Bahrain. Med Bull. 2005;27(4):1–5.
20. Bonvalot S, Desai A, Coppola S, Le Pichoux C, Terrier P, Démont J, Le Cesne A. The treatment of desmoid tumors: A stepwise clinical approach. Annals of Oncology. 2012;23(10):x158-x166.
21. Gluck I, Griffith KA, Biermann JS, Feng FY, Lucas DR, Ben-Josef E. Role of radiotherapy in the management of desmoid tumors. Int J Radiat Oncol Biol Phys. 2011;80(3):787-92.
22. Thomas VT, Jose S. Bilateral breast fibromatosis: Case report and review of the literature. Journal of Surgical Education. 2011;68:320-325.
23. El Demellawy D, Herath C, Khalil M, et al. Fibromatosis of the male breast with concurrent florid-type gynecomastia: Report of a case and review of literature. Pathology—Research and Practice. 2011;207:306-309.
24. de Bree E, Keus R, Melissas J, Tsiftsis D, van Coevorden F. Desmoid tumors: Need for an individualized approach. Expert Rev Anticancer Ther. 2009;9(4):525–535.

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