Case Report

Radiologic presentation of a myofibroblastoma of the adult male breast

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\textbf{Abstract}

We present the case of a 50-year-old male with bilateral gynecomastia who was incidentally found to have 0.8-cm subareolar mass on computed tomography. Mammographic and sonographic characteristics of the lesion are described as well as a brief historical review of myofibroblastoma, a rare mesenchymal tumor.

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\textbf{Introduction}

Myofibroblastomas are rare benign mesenchymal neoplasms composed of stromal fibroblast cells within whorls of collagen. Myofibroblastomas were first described in 1987 as benign breast tumors contained in a pseudocapsule with spindle cells and abundant collagen [1]. These cells show smooth muscle differentiation and stain positive for muscle-specific actin, alpha-smooth muscle actin, and desmin. These tumors are entirely benign and will grow slowly over time if not excised [2]. Myofibroblastomas are most commonly described in the middle-aged male breast, with multiple cases reported in postmenopausal women. There is 1 case report of a breast myofibroblastoma in a 10-month-old male [3]. Though myofibroblastomas are most commonly found in the breast, tumors have been reported in many locations including the liver [4], inguinal canal, posterior vaginal wall, buttoc, anterior abdominal wall, mid back [5], and in the scrotum [6,7]. Some have theorized that these tumors arise from ectopic breast tissue along the embryonic milk lines; however, tumors occurring in regions outside the milk line such as buttoc or liver contradict this theory [5].
Case

A 50-year-old male with a past history of bilateral gynecomastia and sarcoidosis maintained on low-dose corticosteroids for 2 decades presented to the Emergency Department with a chief complaint of hemoptysis. The hemoptysis resolved spontaneously without an identifiable cause. Computed tomography (CT) chest with intravenous contrast was performed to evaluate patient symptoms and revealed an incidental 0.6-cm soft-tissue density mass in the left breast (Fig. 1). Upon further questioning, the patient reported a year long history of a palpable tender mass in the left breast. The patient was referred to breast clinic for further evaluation of this finding.

Imaging and diagnosis

Mammogram confirmed the presence of a circumscribed subareolar mass corresponding to a palpable mass and CT finding (Figs 2A and B). No associated calcifications or architectural distortion were seen. Ultrasound confirmed the presence of circumscribed, nearly isoechoic 0.8-cm mass with minimal internal vascularity on color Doppler assessment without posterior shadowing (Figs 3A and B). There was no axillary lymphadenopathy. An ultrasound-guided core-needle biopsy of the mass was performed, showing immunohistochemical stains positive for vimentin, CD34, CD99, bcl-2, and desmin and negative for AE1/3 consistent with a myofibroblastoma. Surgical resection followed with final pathology confirming a mammillary myofibroblastoma composed of bland-appearing, plump spindle cells in vague fascicular groups with interspersed hyalinized collagen bundles.

Discussion

Myofibroblastomas have been characterized in multiple case reports with a variety of radiographic findings. Typical masses measure 1-4 cm [1,8], with rare cases demonstrating much larger lesions up to 16 cm [9,10]. This case demonstrates a small myofibroblastoma, possibly due to the
workup being trigged by incidental discovery on CT rather than the patient presenting with a larger mass. Mammography characterizes masses as round to oval shaped. They may demonstrate microlobulated margins and are without calcifications or other suspicious features [11–14], similar to the findings in the presented case. Sonographic findings vary from homogeneously hypoechoic to heterogeneous mass with or without posterior shadowing [8,12,14–16]. Doppler may demonstrate variable internal vascularity [9,13]. Sonography in this case demonstrated a homogeneous nearly isoechoic mass with minimal vascularity on Doppler. CT is less often performed but typically shows a circumscribed mass, with 1 case report showing low-density attenuation due to the presence of fat [4]. MRI findings show T1 hypointensity to isointensity with positive early enhancement and nonenhancing septations. The masses are typically T2 hyperintense [4,9,11,13]. Myofibroblastoma is benign with no known cases of lymph node or distant metastasis. The differential diagnosis for a breast mass in a male includes breast cancer, gynecomastia, subareolar abscess, fat necrosis, epidermoid cyst, lipoma, and malignant melanoma [17].

The benign appearance of male myofibroblastoma must be contrasted to the appearance of breast cancer in men, the most severe diagnosis on the differential list for a male breast mass. Mammographic appearance of male breast cancer typically parallels that of a female breast cancer showing an irregular mass with spiculated margins with or without microcalcifications or rarely microcalcifications alone. The lesion size can vary, with a mean size of 2.2 cm ranging as high as 19 cm, with increasing size indicating more advanced disease. Sonographic appearance of male breast carcinoma is most frequently a solid mass with irregular margins and variable vascularity on color Doppler evaluation. Axillary lymphadenopathy occurs in a minority of patients [18]. Although mammographic and sonographic features of myofibroblastoma are not pathognomonic, they are significantly different from the typical appearance of breast cancer and should be considered in the differential list of benign though uncommon masses in the male breast.

Conclusion
The case of a 50-year-old male with a small myofibroblastoma is presented, with imaging findings demonstrating typical appearance on CT, mammography, and ultrasound. Myofibroblastomas represent a rare benign mesenchymal neoplasm that may be encountered in male patients presenting for evaluation of palpable abnormalities in the breast.

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