A Case Report of a Large Myoid Hamartoma of Breast

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Abstract
Breast hamartomas are uncommon breast tumours with unclear pathogenesis & varied histological patterns resulting in under-diagnosis. Immune-histochemistry is a valuable tool in their diagnosis. We report the case of a large myoid hamartoma of breast. Based on our literature search this is the largest reported size of a myoid breast hamartoma that was successfully managed by enucleation.

Keywords: myoid hamartomas, immunohistochemistry.

Introduction
Breast hamartoma is an uncommon breast tumour that accounts for approximately 4.8% of all benign breast masses[1]. It contains lobular breast tissue involving various fibrous, fibrocystic and adipose tissues. The pathogenesis of hamartomas remains unclear and its diagnosis is underestimated by clinicians and pathologists. We report a case of an uncommon myoid hamartoma of breast.

Case Presentation
A 33 year old patient presented with a painless lump in left breast since 10 years, gradually increasing in size. There was no history of nipple retraction or nipple discharge. On examination she had a 15 x 10 x 10 cm mobile well defined lump in the left breast, with smooth surface & soft consistency. There was no palpable axillary, cervical or supra-clavicular lymphadenopathy & the contralateral breast was unremarkable.

Biopsy of the lesion revealed hypo cellular lesion composed of bland spindle to epitheloid cells with abundant pink cytoplasm arranged in solid & nest like patterns. Immunohistochemistry suggested smooth muscle proliferation (SMA & desmin strongly positive & weak expression of CD 34). Intra-operatively the lump was found to be well circumscribed & was enucleated.
Final histopathology report revealed a well circumscribed lesion showing admixture of fibrous tissue, mature adipocytes, numerous smooth muscle bundles & entrapped disordered lobules & ducts suggestive of myoid hamartoma. H&E section at 5X, showing a circumscribed mass composed of adipose tissue smooth muscle and uniform ducts.

IHC- Smooth muscle seen (10x)

Discussion
Breast hamartomas are poorly defined, uncommon, benign breast neoplasms. The average age of the patients with breast hamartoma ranges between 33.5 and 66.5 years. They were initially defined as mastomas in 1928 by Prym. They are well circumscribed, mobile tumours not adherent to skin or muscle. Mammographic studies show a peripheral, lucent halo, and the mass is radiographically similar in pattern to a normal breast (“breast within a breast”). Calcification is absent. USG evaluates breast hamartomas as having sono-lucent fat and a heterogeneous internal echo pattern with echogenic fibrous components. MRI characteristically shows a smooth and well-defined hypointense rim, internal heterogeneous enhancement and the presence of fat density. Histologically, these encapsulated lesions are composed of various breast tissue components such as ducts, lobules, stroma prominently adipose, and smooth muscle, without any particular arrangement. The lobular distribution and the presence of fat in breast hamartomas are distinguishing features compared with fibroadenomas. A fibrotic stroma surrounding the lobules and extending into the interlobular areas, causing obliteration, is the most frequently observed feature & is often referred to as interlobular fibrosis. Hamartoma is referred to as myoid hamartoma, a rarer form, when it shows a significant smooth muscle component.
This term was first used by Davies and Riddell\textsuperscript{[14]} in 1973. Myoid hamartomas can reach large sizes.
The sizes reported in a literature review range from 1.1 cm to 17 cm\textsuperscript{[15]}.
The etiopathogenesis of breast hamartomas is not clear but they are thought to result from dysgenesis rather than a true tumorous process. However, female sex steroid hormones\textsuperscript{[7]} have been implicated in the development of breast hamartomas as demonstrated by estrogen receptor (ER) and progesterone receptor (PR) positivity in epithelial cells and stromal cells\textsuperscript{[6]}. Additionally, there are no clear data on the source of smooth muscle for myoid hamartomas, but this muscle could derive from vessels, the nipples, undifferentiated breast stromal tissue or myoepithelial cells\textsuperscript{[16,17]}. Another hypothesized smooth muscle source is the metaplasia of breast stromal cells\textsuperscript{[17]} into smooth muscle cells. The existence of CD34 on smooth muscles\textsuperscript{[7]} is an important sign of the metaplasia of stromal cells into smooth muscle cells. Myoid hamartomas stain strongly positive for SMA, desmin and vimentin by immunohistochemical staining.

The treatment of these lesions is surgical excision.

**Conclusion**

In conclusion, we present a case of a large myoid hamartoma, an uncommon benign breast tumour. Although breast hamartomas are rare, benign lesions, these lesions can reach large sizes. Based on our literature search this is the largest reported myoid hamartoma. A diagnosis can be made by core needle biopsy & immunohistochemistry, along with an appropriate correlation of clinical and radiologic features. Breast hamartomas may be underdiagnosed because pathologists may categorize these lesions as fibroadenomas instead of hamartomas. Thus, the true incidence may be higher than the literature indicates. With more widespread use of newer imaging modalities & availability of immunohistochemistry, more number of benign breast lump may be accurately identified as breast hamartomas.

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**Conflicts of interest-** none

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