Recurrent Giant Phyllodes Tumour of the Breast
Pathological Considerations and Management Approach for Recurrence and Metastasis*

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ABSTRACT
Phyllodes tumours are rare and account for 0.4% of all breast tumours. The majority of them tend to be less than 5 cms in size with giant tumours larger than 10 cms being about 20% of these cases. They display a broad range of clinical and pathological behavior and are regarded as falling within the spectrum of fibroepithelial neoplasms. Surgery has been the primary modality of treatment. However the extent of resection and the role of adjuvant radiotherapy and chemotherapy are still controversial. The risk of recurrence is (4.7% - 30%) for benign phyllodes tumour and (30% - 65%) for borderline and malignant phyllodes tumour. A case of giant phyllodes tumour measuring 36 × 30 cm and weighing 6.8 kg is presented, which had recurred following a previous resection 5 years back. The literature is reviewed with regards to pathology, the role of investigations, nature of resection and the risk factors for recurrence and metastasis.

Keywords: Phyllodes Tumour, Mastectomy, Giant Tumour, Recurrence

1. Introduction
Phyllodes tumour (PT) are rare fibroepithelial breast tumours that usually occur in adults and are usually seen in women aged 35 to 55 years with only a few cases reported in men [1-4]. They are composed of benign epithelial component and a cellular spindle cell stroma forming a leaf like structure [1-5]. No one morphology is reliable in predicting the clinical behavior of the tumour. While the surgical management of phyllodes tumour has been addressed many times in the literature controversy still exist [1-9]. There are only a few reports however that has specifically commented on giant phyllodes tumour as an entity, which presents to the surgeon with general unique management problem [6]. The median size of the tumour is around 4 to 7 cm [1-8]. Twenty percent of tumours grow larger than 10 cm, the arbitrary cut off point for the defining giant tumours [1-8]. While presenting one of the large giant phyllodes tumour reported in the English literature, the literature is also reviewed with regards to its pathology, management and risk factors for recurrence and metastasis.

2. Case Report
A 62-year-old lady presented with large right breast lump of 14 months duration. This was associated with breast pain. The lump was measuring 36 × 32 cm with an overlying scar of previous surgery being apparent (Figures 1 and 2). The lump was firm and mobile and there were no axillary lymphadenopathy. Five years back she had undergone wide excision of a breast lump measuring 3 × 4 cm in another hospital, which was reported as benign PT with clear resected margin. A CT scan of the present breast lump revealed a large well-circumscribed lobulated mass involving the whole breast with stretching of the overlying skin. A CT scan of the abdomen and chest...
however revealed no metastasis. A core biopsy of the lesion confirmed borderline phyllodes tumour. She underwent mastectomy without reconstructive surgery. The histopathology report confirmed borderline phyllodes tumour (Figure 3) with tumour free resected margins. When last followed up 5 years later she was doing well with no recurrence.

3. Discussion

PT are fibroepithelial neoplasms with epithelial and cellular stromal components, the latter of which represents the neoplastic process [4,5]. The presence of an epithelial component differentiates PT from other stromal tumours [3-5]. They make up to 0.3% to 0.5% of the female breast tumours and have an incidence of about 2.1 per million, the peak of which occur in women aged 45 to 49 years, 15 to 20 years later than that for fibroadenoma [1-8]. The tumour is rarely found in adolescent and elderly. PT usually presents as a rapidly growing but clinically benign breast lump. In some patients, a lump may be apparent for several years, but they only come for evaluation when the mass has suddenly increased in size. Delay between the first symptom and presentation can vary from 2 days to 15 years and a breast lump is noted in 74% of these patients [4]. The other symptoms include mastodynia (14% to 42%) and less frequently skin ulceration (2.4% to 10%) of the cases, which are usually associated with large or malignant PT [4,7]. Due to the large size of the tumour the overlying skin is tense and presents with visible and dilated venous plexus that may eventually ulcerate [1]. The tumour size is reported to vary from 5 to 450 mm with a mean size of 40 to 70 mm [1-8]. PT is unilateral in 96% of cases with very few cases of bilateralism reported [3,4]. They are commonly found in upper quadrant (46%) with an equal propensity to occur in either breast (48.8% to 50%) [3,8].

PT can be extremely difficult to differentiate from fibroadenoma, which is sometimes treated with a non-operative approach. For this reason, early diagnosis of PT, which often includes surgery, is pursued [1-4]. The preoperative diagnosis is reported to be achieved in 10 to 29% of cases using various diagnostic modalities [9]. Ultrasound examination of the breast has distinct advantage as an initial investigation in view of its ease of performing and availability. Various diagnostic features, which are characteristic, include a well-circumscribed lobulated mass, heterogeneous internal echo pattern and lack of microcalcification [4,10]. Patients with fibroadenoma are generally younger than patients with PT and have a larger ratio of length to anteroposterior diameter and are generally smaller in size than PT [10]. Attempt to establish a preoperative diagnosis and to differentiate between malignant and benign PT through the use of colour Doppler ultrasound has been made [11]. The feature that is suggestive of malignancy includes marked hypoechochogenicity, posterior acoustic shadowing and higher vmax (systolic peak flow velocity) [11]. Mammography however is reported to be extremely low in accuracy in differentiating a PT and fibroadenoma. However, in tumours which are large, presence of irregular margin are more likely to be predictive of malignant PT rather than benign PT. MRI has also played a significant role in diagnosing difficult cases. The feature characterizing benign PT included a lobulated or polygonal shaped mass with smooth border, which may be cystic and septated and has a gradual or rapid pattern of time signal intensity curve [12]. The reported pathognomic feature of PT also include characteristic leafy internal morphology, best shown in subtraction MRI which highlights the enhancing cotyledons solid pattern within irregular blood filled cystic spaces [12].

Preoperative histological diagnosis can be achieved by fine needle aspiration (FNA) or core biopsy [10-13]. The role of FNA in establishing a preoperative diagnosis is
limited with reported success of around 12%, the limiting factor being the common cytological feature with fibroadenoma (FA). However, the presence of elongated spindle cells in the background is reported more likely to be present in PT, than FA [13]. Core biopsy is preferred for establishing a preoperative diagnosis, as the histological gain from this procedure is important in guiding surgical treatment. Most of the PT is benign with 15% to 30% being classified as malignant [1-8]. PT is generally divided into benign, borderline and malignant histotype based on the macroscopic appearance of the stromal component [3-5]. A benign tumor is characterized by 0 - 4 mitosis/10 high power field (HPF), predominately pushing margin, and usually 1+ but occasionally 2+ stromal atypia. Borderline tumour are determined by 5 - 9 mitosis/10 HPF, pushing or infiltrating margin, and 2+ stromal atypia and malignant PT is characterized by 10 or more mitosis/10 HPF, predominately infiltrating margin usually 3+ but occasionally 2+ stromal atypia [3-5]. Histological appearance may not however correlate with clinical behavior as both malignant and borderline tumor has been shown to be capable of metastasizing. Histotype however is reported to be an independent prognostic factor with 5 year survival of 95.7% for benign tumors, 73.7% for borderline and 66% for malignant tumours [1,4,7,8]. The 10-year survival drops to 79% for benign and 42% for malignant tumors [1-4,7,8,14-16]. Metastasis has been noted in 25% to 31% of malignant PT but in only 4% of all PT [1-4]. The most common sites of metastasis include lungs; bone liver and distant lymph nodes [1-5,14-16]. Skin involvement with tumor does not appear to be a predictor of metastasis. About 20% of patients have palpable axillary lymphnodes on presentation but only 5% of these show histological evidence of metastasis upon pathological examination [1-5,14-18]. Among the malignant PT only 15% metastasize to axilla [1-4]. The metastatic lesion contains only stromal elements with no malignant epithelial elements observed. In view of the rarity of lymph nodal involvement most of the authors are of the view that routine removal of lymph nodes is not necessary unless they are palpable [1-4].

About 20% of PT is considered giant when the size exceeds 10 cm in diameter [5,14-18]. However the importance of this cut off value has been disputed as there is a continuing debate over the prognostic significance of the tumour size [1,2]. Even though malignant tumours tend to be larger than benign, the malignant histotype correlating with size has not been well established [1,2]. Various factors have been found to be related to clinical behavior and outcome. Of these stromal overgrowth, tumour necrosis and mitotic activity have been found to be the most consistently associated with recurrence, metastasis and poor survival [2,14,15]. Other factors that may correlate with outcome are tumour size [2,3,15,16], stromal atypia [14] and stromal cellularity [17], lymph node metastasis [16], age [2,16], nulliparity and absence of cysts.

Local recurrence usually occurs within few years after the surgery and histologically resembles the original tumor [1-6] [Table 1]. Occasionally recurrent tumour shows increased cellularity and more aggressive histological features than the original lesion [2-6]. This was evident in our case where an initially benign tumour recurred as borderline malignant tumour. In most patients local recurrence is isolated and is not associated with development of distant metastasis [1-4,14-17]. Among these various pathological factors, mitotic activity and inadequate margin clearance following resection was strongly associated with recurrence while stromal cellularity, stromal overgrowth, stromal atypia, along with mitotic activity, tumour margin positivity and heterologous stromal elements significantly correlated with metastasis [3]. All PT can recur regardless of their histotype although the risk of recurrence is low in benign form, ranging from 4.7% to 30%, and notably higher in borderline and malignant form (30% to 65%) [1-5,14-16]. However review of literature of recent publications revealed that local recurrence after margin negative breast conserving resection of borderline malignant and malignant PT was found to be 21.7% and 16% respectively [3,4,19-31] (Table 1) Local recurrence can usually be controlled by further wide excision and mastectomy is invariably not required. Mastectomy is however considered for local recurrence after local surgery has been performed for borderline or malignant tumour [1,2]. Occasionally aggressive local recurrence can result in widespread chest wall invasion with direct invasion of the underlying lung parenchyma. Isolated reports of good palliation in this situation with radiotherapy have been reported [1].

Surgery is the treatment of choice for PT, but the extent of surgery remains controversial particularly for borderline and malignant PT [1-8,14-17]. Most authors agree that the appropriate procedure for benign PT is a wide local excision allowing 1 to 2 cm margin in all directions [1-8,14-18]; however enucleation alone is considered insufficient [15,17]. In the event, the diagnosis of benign PT is a histological surprise following excision of breast lump, re-exploration with wider resection of healthy tissue is not considered justified because of the low recurrence rate [2-4]. While these patients may enter into a surveillance programme, patients with histologically positive resection margin should undergo further surgery [2-4,8]. However in malignant PT and tumours larger than 5 cms, mastectomy is often recommended [2-4,14-16]. Routine resection of the axillary lymph node is not recommended as metastasis is seen in only in around 5% to 10% [1-4,6-8,14-17]. The role of adjuvant radio-
therapy or chemotherapy remains uncertain [8]. Some investigators report that radiotherapy improves disease free survival [32,33]. A significant reduction in local recurrence has been reported when margin negative resection has been combined with adjuvant radiotherapy; Adjuvant radiotherapy following margin negative resection of borderline and malignant PT, lead to reduction of recurrence from 86% without adjuvant radiotherapy to 59% when radiotherapy was used (p = 0.02) [33]. There are others who recommend adjuvant radiotherapy only in cases of incomplete resection [8,32]. Chemotherapy may be considered in patients with stromal overgrowth [15]. However to date neither adjuvant radiotherapy or chemotherapy are routinely recommended [1-4].

4. Conclusion

PT represents a heterogeneous group of tumours sharing the same macroscopic features with an unpredictable outcome. The prognosis depends on the histological and biological characteristics of the tumour rather than their clinical behavior. Giant PT constitutes 20% of these tumours and would invariably need mastectomy. The data concerning the prognostic factors of PT are conflicting and both benign and malignant PT could recur or metastasize. While ultrasound, CT and MRI of the breast could be suggestive of PT, establishing a definite preoperative diagnosis may be difficult and would invariably need core biopsy. Surgery remains the cornerstone of treatment consisting of resection with healthy surrounding breast tissue at initial surgery. Local recurrence can be controlled with further surgery and may not be associated with distant metastasis. Routine axillary lymph node dissection is not recommended. High risk of local recurrence and or metastasis is usually associated in patients with large tumours, infiltrating tumour margin, severe stromal overgrowth, atypia and cellularity.

REFERENCES


