Huge borderline phyllodes breast tumor with repeated recurrences and progression toward more malignant phenotype: a case report and literature review

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Background: Phyllodes tumor (PT) is a rare breast fibroepithelial biphasic tumor composed of stromal and epithelial components. The patients suffering from this disease present with a large, round, mobile, fast-growing lump, and the giant PT of more than 10 cm in diameter is so uncommon. Surgery is regarded as the primary treatment, but curative efficiency of adjuvant chemotherapy and radiotherapy is so indefinite.

Case presentation: We reported one case of a middle-aged woman with a huge borderline PT in the right breast, over 20 cm in size. The pathology of needle core biopsy of the lump was suggestive of PT of the borderline subgroup, and then she underwent mastectomy of the right breast. The patient had recovered well without any postoperative treatment until a local recurrence occurred 1 year after operation. The tumor was removed with lumpectomy, which was pathologically diagnosed as malignant PT. We followed up her by telephone and heard about her postoperative adjuvant radiotherapy and chemotherapy, as well as her well recovery.

Conclusion: The pathology of PT with low incidence is mostly benign, but local recurrence is common, and the histopathology progresses toward worsen trend. Besides, due to the difficulty in precise diagnosis of the borderline PTs, it is recommended that this subtype of patients should undergo total mastectomy. Although the curative effect of postoperative treatment has not been recognized internationally, patients, especially those with huge tumors, may benefit from these treatments.

Keywords: phyllodes tumor, breast neoplasms, recurrence, diagnosis, treatment

Introduction

Breast phyllodes tumors (PTs), composed of stromal and epithelial elements, are rare biphasic fibroepithelial tumors, with an incidence of about 2.1 per million women, accounting for 0.3%–1% of breast neoplasms. Breast PTs, first named by Johannes Muller, have numerous synonyms, such as cystosarcoma phyllodes, adenomatous myxoma, and pseudosarcoma adenoma. WHO decided to use the term “phyllodes tumor” to name this disease formally in 2003. It usually occurs in middle-aged women mostly (age, 35–55 years). Clinically, the variable tumor size is 4–7 cm on average, but about one-fifth of the PTs are called giant PT due to the uncommon diameter of more than 10 cm. Depending on the histopathological features, PTs are categorized into three grades with different proportions, benign (60%–75%), borderline (13%–26%), and malignant (10%–20%). Patients suffering from PTs have no specific clinical manifestations, and it is difficult to distinguish benign subgroup from borderline and malignant subgroups. Lymph node metastasis is rare, and the metastatic path mainly
relies on the blood. Surgical removal is the primary treatment for PT, given that adjuvant treatments play a poorly efficient role. Here, we report an uncommon case of a female patient with a huge breast borderline PT and review literature about PTs.

**Case report**

In April 2016, a 44-year-old premenopausal woman was admitted to the Breast Surgery Department, the Second Hospital of Dalian Medical University, with a 12-month history of a huge lump in the right breast. Initially, the patient just had a small fist-sized lump in the right breast, without any pain or discomfort. It grew so slowly that she did not pay much attention on it. Then, the lump started to grow rapidly and became very large within 2 months, and her right breast was entirely covered with a 8-inch ball-sized mass. The breast skin appeared dark and had ulceration.

The patient had undergone breast lump resection twice in 2001 and 2008, and both pathological diagnoses were benign PT. The patient had no history of cardiovascular or respiratory disease. Moreover, she had no family history of breast cancer.

Physical examination revealed that the left breast was normal, without any palpable lump. The right breast was obviously large, nearly 25×25 cm in size, with a circumference of nearly 80 cm. The skin on the surface of the lump was dark, with ulceration area of 7×5 cm and without bleeding or discharge (Figure 1A and B). On palpation, the size of the lump was 18×20 cm, hard with local tenderness and poor mobility. Axillary lymph nodes were not palpable.

Mammography showed a huge right breast, about 23×23 cm, with dermal ulceration around nipple (Figure 2A and B). Ultrasonoscopy showed the right mammary gland that was enlarged obviously, with inset showing echo intensity disorder and blood flow signal probed (Figure 2C). Several hypoechoic focuses were seen on the area of right axilla, of which the largest was 1.5×0.9 cm in size. The left breast and axilla had no abnormal signal. Biochemical examination revealed no significant abnormalities. The chest computed tomography (CT) revealed that the lump had not broken into the chest cavity, and the lungs were normal (Figure 2D). The patient’s other relevant examinations such as the abdomen CT, the brain CT, and the bone ECT were also normal.

Needle core biopsy of the breast lump was suggestive of PT of the borderline subgroup. Then, the patient underwent mastectomy of the right breast (Figure 3) and sentinel lymph node biopsy. During surgery, surgeons detected that the pectoralis major was invaded, and so the partial muscle adhered to the tumor was resected. The intraoperative frozen section showed no metastatic cancer cell in sentinel lymph nodes. Finally, the latissimus dorsi muscle flap was graft to remedy for the tissular defect on the chest. Postoperative treatment was uneventful, and she was discharged after 6 days.

The cut surface demonstrated a leaf-like pattern. The scope of dermatic ulceration or necrosis was 13×10 cm. Postoperative paraffin-based histopathology showed fast cellular proliferation, mitosis about 9/10 high-power fields (HPFs) (Figure 4A and B). The tumor did not invade the skin. Immunohistochemistry (IHC) was as follows: spindle cells AE1/AE3, smooth muscle actin (SMA) (+), Desmin (−), CD34 partial (+), BCL-2 (−), p53 partial (+), ki-67 15%, tumor cells, estrogen receptor (ER) <1% weakly positive, progesterone receptor (PR) <1% weakly positive, and Ki67 40% (Figure 4C–J).

![Figure 1](https://www.dovepress.com/)

**Figure 1** A huge borderline phylloides tumor in the right breast of a 44-year-old woman: front image (A) and lateral image (B).
The patient had recovered well after the surgery until March 2017. The patient came to our hospital again because of a local recurrence. Her palpation and imaging examinations indicated three large masses on the right chest, the largest one of about 6.7×4.0 cm (Figure 5A–C). Then, she had lumpectomy, and the tumor was diagnosed as malignant PT (Figure 5D). After the second surgery, we followed up her by telephone, and knew that she had underwent radiotherapy and chemotherapy. Up until now (February 2018), she is still alive and undergoing chemotherapy.

**Discussion**

Currently, although a lot of research studies focused on breast PTs, mechanisms and causes of PTs are still far from being clarified. Tan et al\(^8\) reported that the mutation in MED12, RARA, FLNA, SETD2, and KMT2D was observed in PT, suggesting that they may play important roles in PTs tumorigenesis and development. One study by Sapino et al\(^9\) showed that ER-β with high expression in elderly patients was considered as an independent factor, and pregnancy status and a high level of estrogen were also related to this disease. Gong et al\(^10\) studied biological mechanisms of malignant transformation, concluding that miR-21 may drive the progression, and α-SMA and miR-21, as independent prognostic markers, had more predictive value than histologic classification. Hodges et al\(^11\) suggested that some tumors showed the characteristics of both fibroadenoma and PTs, and the deletion or amplification of some allele led to the
transformation of fibroadenoma into PTs. Another research study reported a higher risk of PTs in unmarried, infertile, and postmenopausal women, as well as in women using oral contraceptives at childbearing age.\textsuperscript{12}

Patients suffering from breast PTs usually have non-distinctive symptoms at the beginning, and long medical history ranged from several months to years.\textsuperscript{13} Yet, only a small number of patients consulted doctors because of rapidly increasing tumors. The tumor usually exists in a unilateral breast with the characteristics of hard, clear boundary, oval or phyllodes shape, good tumor mobility, and non-adherence to skin. Huge mass may cause ulcerated, thinned, and tightened skin. The lymph node is involved rarely, and the most common metastasis path is through hematogenous channels, mainly to the lung, the pleura, and the bone.\textsuperscript{14} The tumor with distant metastasis is usually characterized by large mass, invasive growth, obvious hyperplasia of interstitial element, and abundant mitotic activities.\textsuperscript{15}

Due to the lack of typical clinical symptoms and pathological characteristics that can distinguish fibroadenoma, the misdiagnosis rate is high. To diagnose the PTs accurately, it is necessary to find both epithelium and stoma in tissue specimens.\textsuperscript{16} The stromal cell components affect the pathological behavior of PTs. WHO classification of breast tumor in 2012 recommended that the PTs should be classified into three subgroups based on the stromal cellularity, stromal atypia, stromal overgrowth, mitosis, and tumor margins.\textsuperscript{17} Histologically, benign PTs have consistent nuclear morphometry of interstitial nucleus with less than 5 mitoses per 10 HPFs. In some areas where interstitial cells are sparse, hyalinosis or myxoid degeneration can be observed. Malignant PTs reveal the characteristics of nuclear pleomorphism, excessive growth of interstitial cells, and invasive margins, with mitosis more than 10/10 HPFs. Sometimes histological serialities make it difficult to exactly define the grade of tumors.

\textbf{Figure 4} The tissue section showing borderline phyllodes tumor with mucoid degeneration and some areas abounded in cell: (A) 100× and (B) 200×. IHC results of (C) AE1/AE3, (D) SMA, (E) CD34, (F) BCL-2, (G) P53, (H) ki67, (I) ER, and (J) PR. \textit{Abbreviations:} IHC, immunohistochemistry; SMA, smooth muscle actin; ER, estrogen receptor; PR, progesterone receptor.
Preoperative diagnosis of breast PT is uneasy. Ultrasound and mammography showed no specific images, just showing large, round, and lobulated masses with clear boundary. In some mammographic images, a bright halo around the tumor can be seen, due to rapidly growing tumor compressing surrounding tissues. Besides, thick calcification spots are more common rather than scattered tiny ones. The existing research studies do not confirm that magnetic resonance imaging (MRI) has advantages in diagnosis compared with the other imaging examinations. However, some studies found diagnostic differences in MRI: the enhancement curve of benign PT was almost ascending type and that of malignant tumor was flat or platform type. The diagnosis of PTs is mainly based on the histopathological examinations. Core needle biopsy, as a crucial method, has been widely reported that the accuracy rate was as high as 99%, and the positive predictive value and negative predictive value were 93% and 83%, respectively. However, about 39% of the patients may receive a false negative result, so the definite diagnosis still depends on the complete excision of the tumor, on account of histological resemblance between PTs and fibroadenoma. Cimino-Mathews et al broke through from immunological
behavior, indicating that P63, P40 and cytokeratins labeled in malignant PTs, not in lower-grade PTs or fibroadenoma. Nevertheless, they still considered that the accurate diagnosis should not only rely on immunoreactivity.

The primary treatment of PT is surgery regardless of any subgroup, including various operation methods such as lumpectomy, wide local excision, and total mastectomy. In the past, surgeons generally chose simple or radical mastectomy for borderline and malignant PTs. The study by Hassouna et al revealed that radical surgery really improved the survival and decreased the rate of local recurrence. In addition, another study by Barth et al showed that the prognosis of wide local excision combined with postoperative radiotherapy was equivalent to that of radical surgery. However, recently, most of the studies failed to verify advantages of extensive surgery, so surgeons proposed conservative surgery. The study by Tan et al indicated that the state of surgical margin was an important prognostic factor. The resection with wide margin could prevent the recurrence effectively. The National Comprehensive Cancer Network recommends that the range of negative margin should be more than 1 cm. Based on the available clinical data, it is generally believed that the range is enough for patients with benign PTs, but those with borderline or malignant tumors should have extensive resection or even the mastectomy. Although there is no consensus for the indication of total mastectomy, some surgeons believe that this method is required for patients older than 50 years, with tumors larger than 5 cm or with other high-risk factors of recurrence or metastasis. Besides, some surgeons deemed that in view of high recurrence rate in malignant subgroup, as long as the histopathological diagnosis was malignance before or during operation, all patients should undergo total mastectomy regardless of the tumor size. On the basis of the reported conclusions and our own experience, we consider that total mastectomy should be feasible for patients who meet the following criteria: benign or borderline tumors more than 8 cm in size, positive resected margin, local recurrence more than three times, or malignant tumors. Lymph node involvement is barely in PTs, so routine axillary lymph node dissection is often unnecessary, unless preoperatively lymph node involvement has been detected pathologically. For those patients with palpable lymph nodes or huge breast mass, sentinel node biopsy or low-grade axillary lymph nodes dissection is recommended.

The curative effect of postoperative comprehensive treatments such as adjuvant chemotherapy and radiotherapy against malignant PT is limited. An early study by Chaney et al revealed that patients suffering from borderline or malignant PTs have been profited from radiotherapy, especially those with tumors of huge size, positive margin, nuclear pleomorphism, and necrosis. Barth et al and Belkačémi et al also confirmed that postoperative radiotherapy could prolong disease-free survival and lower the risk of recurrence. However, because there are no sufficient multicenter large-scale sample studies, the evidence listed above cannot be completely relied on. Several studies showed a contradictive conclusion that postoperative chemotherapy and endocrine therapy had no significant effect on reducing the rate of recurrence or death.

The overall 5-year disease-free survival rate of PT ranged from 78% to 91%. The rate of recurrence is about 10%–40% for all PTs patients, and malignant or borderline patients have higher rate than benign ones. Most recurrence happens within 2 years after surgery; besides, the pathology of recurrence tumors often shows a worse condition. Patients should review at least once every 6 months during the first 2 years and then once a year thereafter. Since the prognosis was closely related to the histopathological subgroup, the extent of primary surgical resection, and medical history of previous fibroadenoma, patients with these risk factors should have more frequent review.

**Conclusion**

This case report particularly presents a huge borderline PT in the right breast and postoperatively local recurrence with worse histopathology. PT is a rare breast disease but recurs easily. The tumor behavior is very difficult to predict. The primary treatment is operation, yet the efficiency of adjuvant treatment is still under investigation. For patients with invasive chest muscles, adjuvant radiotherapy may decrease the risk of local recurrence postoperatively.

**Acknowledgment**

This study was funded by the Natural Science Fund Project of Liaoning province (No 20170540254) and Medical Science Research Project of Dalian (No 1612019).

**Disclosure**

The authors report no conflicts of interest in this work.

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