Primary alveolar rhabdomyosarcoma of breast in a 13 year old female with cardiopulmonary metastases and cytohistological correlation: A case report

Wamala D1,*, Kitayimbwa P1 and Dworak O2

1Department of Pathology, Mulago Hospital and Makerere University, Kampala, Uganda
2Institute of Pathology, Furth Teaching Hospital of University of Erlangen, Furth, Germany

Abstract

Introduction: Rhabdomyosarcoma is the commonest soft tissue sarcoma of childhood. The tumour commonly occurs in the body regions of the head and neck, genitourinary and extremities. Primary rhabdomyosarcoma of the breast is extremely rare and present diagnostic challenges especially in resource limited centers. It is an aggressive tumour with a poor prognosis especially when diagnosed late. Case presentation: We present a case of a 13 year old female with primary rhabdomyosarcoma of breast metastasized to regional axillary lymph nodes, lung and heart. The patient failed to respond to chemotherapy mainly due diagnostic challenges and succumbed to the disease. Conclusion: Breast masses in young patients should be diagnosed early and accurately, and optimal treatment promptly instituted. The incidence of primary rhabdomyosarcoma of breast is increasing in teenagers, and the tumour has a bad prognosis especially in late stage. We think this case will add knowledge and skills required in histological and cytological diagnosis of breast rhabdomyosarcoma.

Keywords: rhabdomyosarcoma; breast tumour; metastatic disease; cardiopulmonary metastases

Introduction

Rhabdomyosarcoma is the commonest soft tissue sarcoma of childhood, accounting for 5% of malignant tumors in this age category [1, 2]. Two histological subtypes are seen in children, embryonal and alveolar. Alveolar rhabdomyosarcoma occur more commonly in adolescents while embryonal subtype is seen more often in younger children [3]. Common sites of primary disease include the head and neck region, genitourinary tract, and extremities [2]. Primary rhabdomyosarcoma of the breast is extremely rare and poses extreme diagnostic challenges especially in low resource countries where ancillary diagnostics tests are not available.

Case presentation

We present a case report of a juvenile primary breast alveolar rhabdomyosarcoma. A 13 year old Ugandan female presented with a right breast tumour which progressed from a small painless lump that developed soon after her menarche at 12 yrs. The patient had no other complaint by then. Her medical and family history was unremarkable. The patient had previously a right breast lumpectomy performed at a rural health facility. Observations made on the biopsy specimen were erroneously suggestive of fibroadenoma. The lump recurred in the same breast 3 months after surgery, this time associated with severe breast pain and right axillary lymphadenopathy clinically staged-T4N1Mx. The patient was referred to our institution one year after the right breast lumpectomy. She came with right breast tumour, associated nipple oedema and satellite areolar tumour nodules (Figure 1). Chest X ray revealed a right homogeneous opacity and a massive right pleural effusion. The pelvic and lumbar spinal X-rays were normal. Breast ultrasound report showed ill-defined heterogeneous solid mass with calcifications in the right breast covering areolar and sub areolar regions. There was another ill-defined echo complex mass in the lateral...
upper quadrant of the right breast measuring (27x16) mm. Doppler echocardiography revealed arrhythmias. Fine needle aspiration cytology of the right breast showed a highly cellular tumour composed of ovoid to round small neoplastic cells with moderate abnormal mitotic figures (Figure 2).

A cytological diagnosis of malignant right breast neoplasm was made, differential of which included poorly differentiated carcinoma, rhabdomyosarcoma, Ewing’s sarcoma and lymphoma.

The patient was referred to the oncology unit where she completed 3 cycles of cyclophosphamide, adriamycin, 5FU, granistatin and bleomycin for breast carcinoma with no improvement. A repeat surgical biopsy was performed. Histological findings included neoplastic cells with round to oval regular nuclei and a monotonous chromatin pattern. The cells formed aggregates interrupted by fibrovascular septae, and within these aggregates where areas of dyshesion suggestive of an alveolar rhabdomyosarcoma. Immunohistochemistry studies revealed that the neoplastic cells were negative for AE/AE3, LCA, S-100. Chromogranin, synaptophysin and TTF-1 but reactive for alpha-actin and desmin (Figures 3 and 4). A poorly differentiated or epitheloid leiomyosarcoma exhibit the same immunostaining profile but not the histomorphology of round small dark monotonous neoplastic cells separated by fibro vascular alveolar like septa seen in this case report. These findings and the histo-morphological pattern supported the diagnosis of Alveolar rhabdomyosarcoma, and excluded small cell carcinoma, lymphoma, Primitive neural ectodermal tumour (PNET) or epithelial neoplasm.

While in hospital, the patient developed respiratory distress with a high WBC count, and died. Autopsy examination revealed a Firm grey white tumor seeding predominantly the right pleura. The tumour was extensively infiltrating the pericardium and diaphragm both of which were markedly thickened. There were 200 ml straw colored right pleural effusion. Tumor involved the entire right lung, mainly infiltrating along the interstitium of the alveolar septae. A number of tumour nodules were seen on the heart (Figure 7) while both gross and histopathological findings were consistent with extensive tumour infiltration of the myocardium (Figure 5).
Rhabdomyosarcoma, malignant tumor of mesenchymal origin is the commonest childhood soft tissue tumor accounting for approximately 3.5% of the cases of cancer among children aged 0 to 14 years and 2% of the cases among adolescents and young adults aged 15 to 19 years [4]. The most common primary sites of rhabdomyosarcoma are the head and neck, genital urinary system and body extremities [2], it rarely presents as a primary tumor in the breast.

In childhood, there are three important histological types: the embryonal type (60%) and the alveolar (20%) [5]. The embryonal rhabdomyosarcoma predominantly seen in age group 0-4 commonly affects the genital urinary system and head and neck tends to have a better prognosis compared to the alveolar type which often develop in the extremities in children over 10 years, resistant to chemotherapy and metastasis early [6].

Primary rhabdomyosarcoma of the breast are extremely rare with previous studies done by the Intergroup Rhabdomyosarcoma Study group reporting only 7(0.2%) out of 3500 cases studied between 1972 and 1992 [7]. Most patients in young age subgroup 13.6 to 16.9 years (mean: 15.4 years), among the seven IRS patients subgroup studied six (86%) had alveolar rhabdomyosarcoma while one (14%) had embryonal rhabdomyosarcoma [7]. 50% of the patients presented with metastasis whereas three patients has previous diagnosis of fibroadenoma. Our patient had two previous erroneous diagnosis of fibroadenoma, the commonest breast tumor in this age group. This probably explains why diagnosis of rhabdomyosarcoma may be delayed. Mammography and ultrasound are utilized to locate the tumor but cannot distinguish it from a fibroadenoma. Hays et al. [7] advises fine needle aspiration cytology on teenage girls with breast tumor greater than 2 cm.

The reactivity of immunohistochemical vimentin and desmin markers correlates with the degree of tumor cell differentiation, as it does in embryogenesis. Thus, only vimentin is present in the cytoplasm of the most primitive cells, and desmin and actin are acquired by developing rhabdomyoblasts. Antibodies against MyoD1 and myogenin are highly specific and sensitive for rhabdomyosarcoma and are currently used as standard antibodies for diagnosis [8]. We did not have MyoD1 and myogenin, so we depended on the characteristic morphological pattern of the tumor in addition to the available immunohistochemistry panel.

There are three major histological sub types of alveolar rhabdomyosarcoma: those with typical features, a solid pattern, and a mixed alveolar and embryonal appearance [3]. This case is an example of a typical alveolar rhabdomyosarcoma with round small dark monotonous neoplastic cells separated by fibro vascular alveolar like septa (Figure 6). Characteristic translocation t(2;13) (q35;q14) between the long arm of chromosome 2 and the long arm of chromosome 13 and translocation t(1;13) (p36;q14) have been identified in Alveolar RMS [9] resulting in PAX3-FKHR or PAX7-FKHR gene fusions respectively present in 18 of 21 alveolar rhabdomyosarcomas, while the embryonal rhabdomyosarcoma often exhibit deletion of the short arm of chromosome [10].

Cytomorphological features of alveolar rhabdomyosarcoma consist of dyshesive round neoplastic cells with scant cytoplasm, anisonucleosis together with numerous mitosis and apoptotic bodies (Figure 4). Differentials include malignant small round cell tumor characterized by small dark round undifferentiated cells. These tumours include Ewing’s sarcoma, synovial sarcoma, hepatoblastoma, granulocytic sarcoma, intrabdominal desmoplastic small round cell tumor, small cell osteogenic sarcoma, small cell carcinoma, Primitive Neural Ectodermal Tumour (PNET), Non Hodgkin’s Lymphoma and Ewing’s Sarcoma. These poorly differentiated tumours are difficult to diagnose basing on morphology alone ancillary test useful in diagnosis and classification are immunohistochemistry and flow cytometry immunophenotyping. fluorescence
in situ hybridization, electron microscopy and reverse transcriptase polymerase chain reaction (RT-PCR) [11]. The alveolar rhabdomyosarcoma can be confirmed by a FISH utilizing a cytospin preparation to evaluate the neoplastic cells for the FKHR gene rearrangement on chromosome 13q14 [12]. Differentials can be excluded using immunocytochemical markers. Our panel included AE/AE3, LCA, S-100. Chromogranin, synaptophysin and TTF-1 Ewing's sarcoma tumour cells tends to be smaller and uniform and tend to have more finely granular and delicate chromatin pattern.

The prognosis of alveolar rhabdomyosarcoma is worse than that of embryonal rhabdomyosarcoma. Our patient presented with bad prognostic parameters namely the tumour site, patient age, histological type and distant metastases to the heart (Figure 7) and lungs (Figure 8).

Figure 7 Secondary tumour nodules from primary breast alveolar rhabdomyosarcoma nodules on the heart.

Figure 8 Secondary tumour nodules from primary breast alveolar rhabdomyosarcoma nodules on the lung

Conclusions

It is important for the medical team involved in management of young people with breast disease to ensure that accurate timely diagnosis and prompt optimal treatment of breast lumps and to be aware of the possibility of rhabdomyosarcoma, an increasingly common sarcoma in children. Fine needle aspiration is increasingly playing a pivotal role in good management of breast lumps especially in teenagers.

Acknowledgements

We are grateful to the patient’s mother who kindly agreed and consented to the publication of this article. We appreciate the support of Institute of Pathology, Frether Hospital for all the immunohistochemistry tests that enabled the accurate diagnosis.

Conflict of interest

All the authors declare that they have no conflict of interest.

References