Phyllodes Tumor of Anogenital Mammary-like Glands with Diffuse Pseudoangiomatous Stromal Hyperplasia

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ABSTRACT

Anogenital mammary-like glands may give rise to various pathologic lesions identical to those known in mammary pathology. Tumor occurring in the anogenital region is extremely rare. The histogenetic origin of this tumor is controversial as it is being debated whether such lesions evolve from ectopic breast tissue and most recently, anogenital mammary-like gland. We report a 28-year-old girl who presented with a painless mass in the anogenital region, which was subsequently excised. Microscopic examination revealed morphologic pattern characteristic of benign phyllodes tumor with pseudoangiomatous stromal hyperplasia. We present this case to emphasize the importance of recognizing this uncommon lesion occurring at an extremely unusual site. We also discuss the histogenesis of phyllodes tumor and related lesions occurring in the anogenital region in light of the current literature along with a brief review of the previously reported cases of anogenital mammary-like glands.

Key Words: Phyllodes tumor, Pseudoangiomatous stromal hyperplasia, Female genital diseases, Perineum

INTRODUCTION

Since the first description by Hartung of mammary tissue that had developed in the vulva of a 30-year-old female in 1872 many neoplastic and reactive conditions that are morphologically very similar to the lesions commonly seen in normal breast tissue have been described in the vulva or anogenital region (1). Such lesions include lactation glands, lactation adenoma, hydrocystoma, hydroadenoma papilliferum, non-mammary Paget disease and invasive adenocarcinoma (2). It is still unclear whether such proliferative processes in the anogenital region originate from ectopic mammary tissue or local adnexial structures. Phyllodes tumor of the breast is a relatively rare fibroepithelial lesion that makes up less than 1% of all primary breast tumors and is very rarely localized at the vulva, or the perianal or anogenital region and only a few cases have been reported to our knowledge (3-7).

We present a 28-year-old woman with benign phyllodes tumor in the perineum with widespread pseudoangiomatosis stromal hyperplasia findings. We also discuss information in the literature on the histogenesis of relevant lesions in the vulva, perineum or anogenital region, their histological features, and previously reported phyllodes tumor cases at similar localizations.

CASE REPORT

The presented case is a 28-year-old patient who presented with a painless mass at the perineal region. The mass had a polypoid appearance and soft consistency. It measured 4.5x3.5x3 cm and was surrounded by a rim of remarkable skin. The cut surface was pale pink-white in color and in a granular fashion. Microscopy of samples showed mammary-like glands under the squamous epithelium and neighboring fibroepithelial tumor with regular borders (Figure 1). The whole tumor was processed and evaluated with multiple serial sections. Similar to the first samples, there was a marked fibroepithelial development pattern with occasional epithelial hyperplasia morphologically (Figure 2). The stroma was only mildly cellular in many areas. Stromal atypia or mitosis were absent (Figure 3). No necrosis was observed. Foci consisting of hyalinized collagen bands and slit-like spacing anastomosing with each other and lined with unspecified cells with no atypia or mitotic activity and no erythrocytes were present in the stroma (Figure 4). This specific hyalinized appearance of the stroma was interpreted as pseudoangiomatous stromal hyperplasia (PASH). There was widespread PASH around the mammary-like glands in the non-tumoral tissue. Widespread ductal hyperplasia was seen in the epithelial component. Widespread apocrine metaplasia and...
hyperplasia were also present. The tumor was quite close to the marked surgical border in some samples.

Immunohistochemical methods revealed CK7 (+), CK20 (-), CEA focal (+), GCDFP-15 focal (+), estrogen receptor 10% (+) and progesterone receptor 90% (+) in the epithelium. The stromal cells and the slits in the PASH areas stained with vimentin, actin and CD-34 (Figure 4). The slits were negative for CD31 and factor VIII.

The lesion was diagnosed as “Benign Phyllodes Tumor” with these morphological findings. Long-term follow-up for recurrence was recommended as the lesion was very close to the surgical border and it can be difficult to predict the prognosis of phyllodes tumors at such rare localizations.

DISCUSSION

Mammary-type tissue in the vulva was thought to be and accepted as ectopic mammary glands representing caudal remnants of the milk line for a long time after its description (1). The presence of ectopic breast tissue neighboring the lesion or in direct histological continuity with the lesion in some of the presented cases has also supported this notion for a long time. However, van der Putte has stated in 1991 and 1994 that this tissue represents the normal structure of the anogenital region and that similar lesions in the perianal region cannot be explained with the milk line theory (1,8). Such gland structures are currently defined as anogenital mammary-like glands. van der Putte has
described the histological features of these lesions. Such glands are larger than apocrine or eccrine glands and have large ducti with acini, diverticuli, short branches and rarely lobules, similar to mammmary glands. Ultrastructurally, the secretory epithelial cells contain electrulcent secretory granules (1,8). Scurry et al have supported van der Putte's theory that secretory epithelial cells contain electrulcent secretory lobules, similar to mammary glands. Ultrastructurally, the large ducti with acini, diverticuli, short branches and rarely glands are larger than apocrine or eccrine glands and have described the histological features of these lesions. Such negative with factor VIII. This immune staining pattern with vimentin, partially positive with smooth muscle actin and mitosis in the lining epithelium. Immunohistochemistry no erythrocytes within the vessel-like slits and atypia or similarity to low-grade angiosarcoma. However, there are of PASH that consists of vessel-like slits is the diagnostic also indicates that the cells lining the vessel-like slits are not of endothelial origin and possibly belong to the spectrum of cells that develop from young mesenchymal cells to myofibroblasts. The fact the cells surrounding the slits were immunoreactive with CD34 and smooth muscle actin and did not stain with endothelial markers other than CD34 in our case supported a myofibroblastic origin and decreased the histopathological possibility of an angiosarcoma.

Malignant or benign, epithelial or stromal lesions of the AGMLG show significant homology with those seen in the breast (12-14). Phyllodes tumor of the AGMLG is a biphasic fibroepithelial neoplasm consisting of an epithelial glandular component and a usually more dominant stromal component. Similar to those in the breast, the definition of a neoplasm as a fibroadenoma or phyllodes tumor is based on the amount and appearance of the stromal component. The incidence of phyllodes tumor affecting the AGMLG is very low and these tumors have generally been reported as single case reports in the literature. The tumor in our case was relatively large and had fibroepithelial morphology with excess stromal development under squamous epithelium with regular borders. The entire tumor was evaluated and the stroma was mildly cellular in many areas but no stromal atypia, mitosis, heterologous differentiation or necrosis was seen. Widespread ductal hyperplasia was seen in the epithelial component, together with widespread apocrine metaplasia and hyperplasia. All features were consistent with benign phyllodes tumor. The differential diagnosis of this tumor includes some stromal tumors such as like fibromatosis, angiomyoﬁbroblastomatosis, angiomyxoma. The epithelial islands may look surrounded by stroma in such mesenchymal tumors but the leaf-like pattern of phyllodes tumor is not present. Papillary hydroadenoma looks like benign phyllodes tumor superficially but there is no stromal component. Two other lesions with biphasic appearance that need to be considered in the differential diagnosis are chondroid syringoma and uterine cervix mullerian adenosarcoma that has metastasized to the anogenital region. However, the characteristic “leaf-like” pattern and biphasic morphology with marked stromal component in phyllodes tumor help in the differential diagnosis with such lesions.

In conclusion, we reported an unusual case of benign phyllodes tumor with diffuse pseudoangiomatous stromal hyperplasia findings in the perineal region to highlight the current concepts regarding the origin of the tumor at the unusual site. We think that the most likely source of phyllodes tumor in anogenital region and the various other proliferative processes that show striking homology.
with lesions occurring in the normal breast is anogenital mammary-like glands. We would like to note that these lesions can be accompanied with the special morphological appearance of pseudoangiomatous stromal hyperplasia, complicating evaluation of the stromal component.

**REFERENCES**


