Angiomatoid Fibrous Histiocytoma: A Rare Entity with Recognized Diagnostic Pitfalls

Anjiomatoid Fibröz Histiositom: Teşhis Tuzaklarına Sahip Nadir bir Antite

INTRODUCTION

Angiomatoid Fibrous Histiocytoma (AFH) is a rare soft tissue tumour. It mainly affects children and young adults and is believed to be of indeterminate differentiation. AFH arises from the dermis and subcutaneous tissue and mostly affects the extremities. It is histologically characterized by thick fibrous pseudo capsule, lymphoid infiltrate with follicular formation and nodules or fascicles of either spindle cells or histiocytoid cells. Pseudoangioectatic cystic spaces filled with blood and lined by tumour cells are the main feature. Immunohistochemistry is usually utilized to aid diagnosis. Classically, AFH is positive for Desmin and CD99 with variable positivity for Calponin, CD68 and EMA. Stains with Factor VIII, CD34, CD31, S100 and keratin are negative (1,2).

The most common genetic abnormality is t(2;22) resulting in EWSR1-CREB1 fusion. Other observed gene fusions are FUS-ATF1 and EWSR1-ATF1.

CASE REPORT

We received case for consultation from a private care provider in Qatar. The case was that of a 10-year-old boy that presented with a painful swelling on the back since 5 months. The swelling was near the dorsal spine, 2cm in size on the right side, not adherent to the underlying structures. The family did not notice any increase in size of the swelling.

Excisional biopsy was done and report from the private care provider suggested soft tissue sarcoma, the differential diagnosis included Ewing/PNET (Primitive neuroectodermal tumor) and Low grade myofibroblastic sarcoma. Accompanying was report from Heidelberg Germany which had contrasting views regarding the diagnosis, which was finally confirmed by Brigham and Women’s hospital, Boston. We review literature and discuss the possible differential diagnosis.

Key words: Angiomatoid fibrous histiocytoma, round cell tumour, soft tissue sarcoma, misdiagnosis, genetic testing
phenotype favouring embryonal rhabdomyosarcoma”. According to Heidelberg Hospital, it was positive for Desmin and focal positive for SMA, CD99, CD34, CD31 and S100. Stains with Pan-cytokeratin, neuroendocrine markers (synaptophysin and chromogranin), CD68 and bcl2 were negative.

We re-evaluated the sections as third opinion and a malignant soft tissue tumour was suggested. The tumour histologically exhibited short, oval spindle cells with overlapped nuclei arranged in plump fascicles and vague storiform growth pattern. Also apparent is fibrous capsule and lymphocytic infiltrate (figure 1).

Brisk mitosis with abnormal figures is present with tumour necrosis. There are also areas of pleomorphic spindle cells in desmoplastic stroma. On immunoprofiling it shows strong positivity for CD99 and Desmin. Staining with EMA showed focal equivocal positivity while AE1/3, CD68 and S100 are negative (figure 2). As per the given picture, rhabdomyosarcoma was the most probable diagnosis.

The patient then travelled to Children Hospital of Philadelphia where a diagnosis of Angiomatoid Fibrous Histiocytoma was made. Interestingly, genetic testing was also done which turned out to be negative for EWSR1-CREB fusion. Wider excision was performed, and it was clear of neoplasia.

Fifth and Final opinion was sought from Brigham and Women’s Hospital Boston, which confirmed the diagnosis of Angiomatoid Fibrous Histiocytoma without further cytologic study.

Currently the patient is on regular follow up and no recurrence has been noted.

DISCUSSION

The problems encountered while dealing with a rare disease are less exposure of the doctor to the disease and the fact that the disease itself has not been studied very much because of its rarity. Thus, with a disease like AFH, which can have a histologic picture similar to sarcomas, the diagnosis and treatment is usually delayed with high possibility of misdiagnosis.

AFH is a rare soft tissue tumour comprising of 0.3% of all soft tissue tumours (1). It arises from the dermis and subcutaneous tissue and mostly affects the extremities. Macroscopically, AFH appears as a lymph node with hemorrhagic area and appears in areas of normal lymphoid tissue such as the antecubital fossa, axilla, inguinal and supraclavicular regions (2). Local symptoms such as pain and tenderness are not common but anemia, pyrexia and weight loss are seen in some cases suggesting release of cytokines from the neoplasm. The median age of presentation is 14 and 20 years as per two studies (3). The metastasis rate is 4% and recurrence rate is that of 12% according to one study which can be attributed to incomplete resection. Other studies report 1% metastasis rate.

AFH is recognized by Pseudoangioectatic cystic spaces filled with blood and lined by tumor cells. There is covering of a thick, fibrous pseudo capsule and inside are nodules or fascicles of either spindle cells or histiocytoid cells. Lymphoid infiltrate with follicular formation is also present in most cases.
However, the characteristic pseudo capsule and lymphocytic infiltrate might either be absent or not present in the representative sample. Fine needle aspiration cytology is thus not recommended (5). The tumor cells are arranged in a variety of growth patterns including sheets, whirls or fascicles. There are also varieties reported with prominent myxoid matrix and schwannoma like features (6-7).

A subset of AFH has a predominantly small blue cell appearance, which can be confused with high grade undifferentiated sarcomas as was present in our patient. This subtype has dark hyperchromatic nuclei with scant eosinophilic cytoplasm mimicking Ewing sarcoma. More pleomorphic examples with brisk mitotic activity have been reported with same prognosis. This subtype is noted in only one report before (8). And it’s highly likely that it is confused for small blue cell tumor especially when degree of suspicion of AFH is low and genetic testing is either not done or turns negative as happened in the case presented.

Immunohistochemistry is usually utilized to aid diagnosis of sarcomas. Classically, AFH is positive for Calponin, CD68, Desmin, EMA, CD99, actin. With Factor VIII, CD34, CD31, S100 and keratin, it stains negative. The most common genetic abnormality is t(2;22) resulting in EWSR1-CREB1 fusion which was observed in eight out of nine tumors analyzed (9). Other observed gene fusions are FUS-ATF1 and EWSR1-ATF1(10).

Rhabdomyosarcoma, on the other hand, accounts for approximately 40% of pediatric soft tissue sarcomas (11). Embryonal rhabdomyosarcoma (ERMS) comprises the single largest category of soft tissue sarcomas in children and adolescents with 4.5 cases per million persons aged 0-20 years in USA (12). It gives a microscopic picture of primitive mesenchymal cells with stellate nuclei with sparse cytoplasm, characteristic “strap”; “tadpole cells” are present in a number of cases. Markers of skeletal muscle differentiation aid diagnosis of ERMS. In alveolar rhabdomyosarcoma, there are aggregates of poorly differentiated round cells that frequently show central loss of cellular cohesion and formation of irregular “alveolar” spaces. The cells in the center are often poorly preserved with degeneration and necrosis. Pleomorphic rhabdomyosarcoma, as another differential to AFH, was previously confused with AFH and is still largely differentiated with the help of immunohistochemistry. Rhabdomyosarcoma, therefore, can easily be mistaken as it also stains positive for CD99 and Desmin.

Extra skeletal Ewing sarcoma/PNET can present similar to AFH with a multilobulated soft swelling in less than 30 yrs. of age patient and hemorrhage, cyst or necrosis on cut surface. On microscopic examination it consists of uniform round cells with round nucleus, distinct nuclear membrane, fine powdery chromatin and one or two nucleoli. It has high vascularity with occasional pseudovascular or pseudoalveolar pattern caused by small fluid filled pools and thus can be mistaken for angiosarcoma, alveolar rhabdomyosarcoma or even AFH. On immunoprofiling Ewing/PNET stains positive for CD99 but not Desmin. Genetic testing for EWS gene fusion help in diagnosis as t(11;22)(q24;q12) is detected in 90% of the cases(13).

Other possible misdiagnoses include Nodular Kaposi with its spindle cell morphology and vascular spaces. To differentiate, the pseudoangiomatoid
spaces in AFH are lined by flattened tumor cells and they stain negative for CD 34 and HHV8. Spindle cell hemangioma typically occurs in the dermis and subcutis of the distal extremities. It contains cavernous vascular spaces, but in contrast to AFH these are lined by an attenuated layer of endothelial cells. The bland ovoid cells of AFH can sometimes be mistaken for granulomas.

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REFERENCES