Follicular dendritic cell sarcoma with paraneoplastic pemphigus: Rare case and a brief review of literature

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INTRODUCTION

Follicular dendritic cell sarcoma (FDCS) is a rare neoplasm (0.4% of soft tissue sarcomas) originating from follicular dendritic cells (FDCs), a non-lymphoid, non-phagocytic antigen presenting cells of the lymphoid system. The tumor was first characterized by Monda et al., and subsequently, specific immunohistochemical markers have been identified. Since that time, only 3 cases of FDCS arising from castleman’s disease (CD) with paraneoplastic pemphigus (PNP) have been reported in the literature.

CASE REPORT

A 20-year-old boy presented with 7 months history of multiple polymorphous vesiculobullous mucocutaneous eruptions around the lips and subsequently in buccal mucosa causing dysphagia. He later developed conjunctivitis associated with blurred vision and atypical targetoid lesions with crusts surrounded by erythema scattered on the trunk and extremities including prepuce and perianal region [Figure 1]. He lost 5 kg during this period. Direct immunofluorescence of biopsy specimen from Skin lesion showed deposition of Immunoglobulin G (+) and C3 in the intercellular region of epidermal cells mainly in the suprabasal region. Linear deposition of IgG is also seen in dermo-epidermal junction suggestive of PNP. Computed tomography (CT) scan revealed a well defined intensely enhancing mass lesion with central necrosis in the right hemipelvis, 6 cm in size, abutting the right obturator internus muscle and displacing the urinary bladder anteriorly. Engorged vessels were seen around the mass [Figure 2]. Exploratory laprotomy was done but the mass was found to be unresectable. Histopathology of the mass showed oval to spindle cells in a background of vascular stroma infiltrated by lymphocytes. The cells were immunopositive for CD21, CD35, CD23, S-100, Claustrin, CD34, CD31, S-100; Leucocyte common antigen, and Human leukocyte antigen-DR were also positive in some cells. Hyaline-vascular follicles were identified at the peripheral portion or occasionally scattered in the center of the tumor. Rich vascularity was highlighted by the CD34 immunostaining [Figures 3 and 4]. Overall features favoured FDCS. The presence of rich vascularity around the tumor and few hyalanized vascular follicles found in histopathological examination gives the clue that the tumor might have developed from Castleman’s disease (CD). As for the mechanisms by which CD induces PNP, it has been proposed that autoantibodies secreted from the Castleman’s tumor play pivotal role. This hypothesis seems to be supported by the present case, in which CD may have triggered both the FDCS and the PNP.

KEY WORDS: Castleman’s disease, follicular dendritic cell sarcoma, paraneoplastic pemphigus

ABSTRACT

Paraneoplastic pemphigus (PNP) is often a fatal autoimmune bullous disease characterized by severe stomatitis, polymorphous skin eruptions, and underlying neoplasms. We describe a patient with PNP associated with follicular dendritic cell sarcoma (FDCS), a rare neoplasm originating from follicular dendritic cells, which are non-lymphoid, non-phagocytic accessory cells of the lymphoid system and play an integral role in regulation of the germinal center reaction and present antigens to B-cells. The presence of rich vascularity around the tumor and few hyalanized vascular follicles found in histopathological examination gives the clue that the tumor might have developed from Castleman’s disease (CD). As for the mechanisms by which CD induces PNP, it has been proposed that autoantibodies secreted from the Castleman’s tumor play pivotal role. This hypothesis seems to be supported by the present case, in which CD may have triggered both the FDCS and the PNP.
FDCS arises from FDCs which are localized within lymphoid follicles; morphologically have entangled cytoplasmic processes and frequent desmosomes; have the ability to trap and retain antigens for long periods of time in the form of immune complexes, but have a non-phagocytic activity; and express molecules involved in the proliferation and differentiation of B-cells.[2] The only predisposing factor identified for FDCS is hyaline-vascular CD (10-20% of cases).[1] FDCS are associated with Epstein barr virus (12%) especially those occurring in the liver.[1] It generally affect young to middle-aged adults, with a mean age of 43 years with no sex predilection.[8] They usually present as painless coalescent nodal mass (60%) of variable size (1-20 cm diameter), but extranodal disease is most commonly seen in the head and neck (tonsil, nasopharynx) and intraabdominal organs (in particular the pancreas and peripancreatic tissues).[3] Other locations: axilla, gastrointestinal tract (17%), liver, oral cavity (13%), mediastinum (8%) and spleen (2%).[7]

Grossly it appears as a firm white mass. HPE showed ovoid to spindled cell whorls with long cellular processes, distinctive nuclei, speckled to vesicular chromatin; sometimes multinucleated cells. Hemorrhage or necrosis is an uncommon finding.[1] Immunohistochemistry positive for CD21, CD23, CD35, Fascin, Clusterin, podoplanin, Desmoplakin Epithelial membrane antigen, Epithelial growth factor receptor, Anti-Follicular Dendritic Cells antibody(Ki-M4), Anti-Follicular Dendritic Cells antibody CNA 42, CD68, Fragment crystallizable region, S100 ± (but not strong) Factor XIIIa ± while negative for Leucocyte common markers (CD1a), cytokeratin, lysozyme, CD34, CD3, CD79a, CD30, Human Melanoma Black HMB45.[2]

A subset of these tumors has been reported to arise in or concomitantly with foci of CD[7] as in our case. The presence of rich vascularity around the tumor and few hyalized vascular follicles found in HPE gave the clue that the tumor might have developed from CD. In fact,
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some argue that FDCS actually develops from CD as a hyperplasia — dysplasia — neoplasia sequence[7] so FDCS may share some of the radiologic features of CD. In our patient, we saw a large lobulated soft-tissue mass with engorged vessels seen around it on CT, the feature of CD.

FDCS behaves more like low-grade sarcoma than lymphoma. Complete surgical resection is considered the treatment of choice; the role of adjuvant therapy (chemotherapy or radiation) is unclear because of the rarity of the disease. Cyclophosphamide, Adriamycin, vincristin, prednisolone CHOP[8] have shown good response. In a small series by Soriano et al.[9] patients who received all three modalities of treatment upfront had the longest disease-free intervals. Sarcoma-based regimen gemcitabine and taxane,[9] imatinib in combination with gemcitabine and cisplatin,[10] rituximab[11] was also reported to have some activity against FDCS. Our patient was given R-CHOP. But response could not be assessed as he expired post first cycle due to pseudomonas infection.

Poor prognostic factors like extensive necrosis, a mass larger than 6 cm, cytotolytic atypia, intraabdominal location, and a high proliferative index increase the likelihood of recurrence, metastasis, and mortality.[7] The lung, liver, and lymph nodes are the most common sites for metastasis, and the tumor may be highly aggressive and fatal within 2 years of diagnosis.[7] Local recurrence rate was reported to be 43% while distant metastases – 24% (lung, liver, peritoneum, pancreas and lymph nodes). Five-year recurrence-free survival rates of 27% to 32%. And overall 5-year survival rate of 79%.[7]

Our patient presented with PNP, an autoimmune bullous disease characterized by severe stomatitis, polymorphous skin eruptions that occurs in association with various forms of underlying neoplasia. It is characterized by the presence of autoantibodies that react with intermediate filament-associated proteins in desmosomes and hemidesmosomes; desmoplakin, bullous pemphigoid antigen 1, and envoplakin.[4] A review of 163 case reports of PNP reported that 84% of cases were associated with hematologic malignancies (Non hodgkin lymphoma [39%], Chronic lymphocytic leukemia [18%], CD [18%], thymoma [6%], Waldenstrom macroglobulinemia [1%], Hodgkin’s lymphoma [1%], and monoclonal gammopathy [1%]). The remaining 16% were associated with nonhematologic neoplasms (epithelial origin carcinoma [9%], mesenchymal origin sarcoma [6%], and melanoma [1%]). Oral ulcerations were the presenting feature in 45% of all cases and appear to be a key feature of PNP.[12]

In conclusion, our case report is unique because it reports for the 1st time to our knowledge as a pathological constellation of follicular dendritic cell sarcoma with castlemean’s disease and paraneoplastic pemphigus arising from extranodal pelvic region, a rare disease at a rare site. Its importance lies in suspicion of diagnosis and considering the treatment of paraneoplastic pemphigus according to the underlying pathology. Being rare its optimal treatment yet to be defined.

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