Infantile Inflammatory Myofibroblastic Tumor with Terminal Ileum Obstruction

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ABSTRACT

Inflammatory myofibroblastic tumor is an uncommon benign neoplasm and its presentation in small bowel is rare. Due to clinical manifestation, laboratory data and radiologic results, there is an increased risk of over diagnosis as malignant tumors particularly malignant lymphoma in childhood therefore may be resulting in wrong therapeutic approach. However pathologic findings could be demonstrated definite diagnosis based on markedly proliferation of bland looking spindle cells which their presence can be confirmed by immunohistochemistry staining to show myofibroblastic differentiation. We describe here an unusual presentation of inflammatory myofibroblastic tumor of the terminal ileum in a 19-month-old girl who presented with anemia, eosinophilia, intestinal obstruction, focal ossification and dystrophic calcification within the tumor.

Keyword: Abdominal Tumour, Inflammation, Myofibroblast

Introduction

Inflammatory myofibroblastic tumor (IMT) is a rare benign lesion. It has a wide range of clinical behavior according to the site of the lesions (1-3). Synonyms that frequently were used in the medical literature are as follows: inflammatory pseudotumor, plasma cell granuloma, plasma cell pseudotumor, xanthomatous pseudotumor, pseudosarcomatous myofibroblastic proliferation, and inflammatory myofibrohistiocytic proliferation (4). The etiology of IMT is still unknown but development of IMT has been described after trauma, surgery or infection (5). There have also been some reports regarding the association of IMT with Behçet’s disease, Hodgkin’s disease or chronic peptic ulcer (6). Although IMT is generally considered to be a benign lesion, there have been reports regarding recurrence, especially in cases of incomplete resection and extrapulmonary lesions, which are large and locally invasive (7). There is a general agreement that the GI tract lesions are predominantly submucosa and composed of inflammatory plasma cells, histiocytes and lymphocytes in a matrix of
spindle-shaped myofibroblasts (8, 9). They are most commonly confused with sarcomas (10). A review of literature for this rare condition was done to delineate the natural history of this tumor and to do a histological confirmation of its benign nature. Because of the risk of local recurrence, IMT cases should have a long-term follow up. We report here this lesion in a 19-month-old girl with unusual clinical presentation of refractory anemia, intestinal obstruction with severe calcification and ossification.

**Case Report**

A 19 month-old girl admitted with abdominal distention. She had a chronic anemia (Hb:5.5 mg/dl) for about 3 months ago which was detected and treated symptomatically. Recent presentation was intermittent abdominal distention after feeding and constipation resulted in acute-abdomen presentation. Other lab data were normal limit in biochemistry, hormone assays but 15% eosinophils in peripheral blood. Abdominal ultrasonography revealed a mass in terminal ileum which was partially obstructed the lumen (Fig. 1).

**Fig. 1:** Ultrasonography showed a mass in terminal ileum which was partially obstructed the lumen.

Total resection of mass was performed and gross examination revealed a well circumscribed firm to hard exo and endophytic creamy-white mass at terminal ileum measuring 9x5x5 cm. Cut surface showed focal bony consistency and whorling appearance (Fig. 2).

**Fig. 2:** A well formed exo-endophytic mass with homogeneous creamy white surface and focal bony consistency at terminal ileum.

H&E sections showed proliferation of spindle cells with oval to spindle shaped nuclei and eosinophilic to amphophilic cytoplasm which were widely separated and haphazardly distributed among with intercellular collagen. In addition there was mixed inflammatory infiltrate composed of myofibroblasts, histiocytes, lymphocytes, plasma cells and rare eosinophils. Areas of severe calcification were present. There was not any mitoses, atypia or nuclear hyperchromasia (Fig. 3).

**Fig. 3-** Proliferation of spindle cells among with intercellular collagen and mixed inflammatory infiltrate with prominent number of eosinophils and plasma cells. Foci of calcification are presented at upper left of the picture (Hematoxylin-Eosin staining, magnification ×400).
Immunohistochemical staining results were as follows: (Fig. 4)

**S100:** Strongly positive reaction in spindle myofibroblastic cells

**SMA:** Strongly positive reaction in spindle myofibroblastic cells

**Vimentin:** Strongly positive reaction in spindle myofibroblastic cells

**ALK1:** Weakly positive reaction in spindle myofibroblastic cells

**CD34:** Negative reaction in spindle myofibroblastic cells

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**Fig. 4 - (A-E).** Immunohistochemical staining. A-C: Showed strongly positive cytoplasmic reaction in spindle myofibroblastic cells in S-100, SMA, Vimentin (magnification ×400). D: Showed weakly positive reaction of nuclei of spindle myofibrablastic cells by ALK1 (magnification ×1000). E: Showed negative cytoplasmic reaction in spindle myofibrablastic cells and positive internal control of vessels (magnification ×400)
Discussion

IMT occurs primarily in children and young adults but can occur at any age (1-3). Mean age of disease is approximately 10 years (11) and infantile cases have also been described (12, 13). Females are affected slightly more commonly than males (14). IMT was originally reported in the lung (15). Extra pulmonary sites including the bronchus, liver, small bowel, mesentery, bladder, kidney, spinal canal, meninges, stomach, pancreas and thyroid have also been reported (16).

Patients with intra-abdominal tumors most commonly presented with abdominal pain and mass with increased girth, or occasionally, with intestinal obstruction (12). Suster reported that some patients might be presented with non-specific systemic symptoms as fever and weight loss. Children may also have growth failure. Common abnormalities presented with laboratory tests include an elevated erythrocyte sedimentation rate (ESR), leukocytosis, thrombocytosis, and hypergammaglobulinemia (17). These symptoms and laboratory abnormalities may disappear after surgical intervention and be distinct from those of sclerosing mesenteritis, usually diagnosed in the absence of constitutional symptoms and laboratory abnormalities (18). In our case anemia might be due to secondary effects of malabsorption of hemoglobin biochemical structure precursors or chronic blood loss of slow growing tumor in terminal ileum. Severe calcification and ossification of this entity due to chronicity of lesion had not been reported previously even in two other neonatal cases with abdominal obstruction (19, 20).

IMT shows three major subtypes in histologic appearance: fibromyxoid and vascular pattern, proliferating pattern, and sclerosing patterns. Immunohistochemistry have revealed that spindle cells of these lesions have originally myofibroblastic differentiation (6). The spindle cells have reaction with antibodies against vimentin, smooth-muscle actin (SMA) and muscle-specific actin. Differential diagnoses of IMT include the gastrointestinal stromal tumor (GIST) and myofibroma for the benign lesion. Myofibroma shows nodular pattern of myoid cells around thin-walled blood vessels. GIST can be excluded by histologic features and positive to CD117 in contrast to IMT. The gastrointestinal autonomic nerve (GAN) tumors are histologically similar to IMT, but they are uniformly negative to muscle-specific actin and CD 68 (21).

Although neoplastic lesions generally have a benign behavior, intra abdominal and retroperitoneal lesions of this type have typically higher local recurrence rates and even more distant metastases (13). Intra-abdominal IMT has a propensity for more aggressive clinical behavior than their extraabdominal counterparts (12). Clonal cytogenetic abnormalities have been demonstrated in some cases (22-24) particularly defined abnormalities on chromosome 2 that result in ALK re-arrangement and is more commonly seen in pediatric than in adult cases (25, 26). Biselli (27) recently found that almost 50% of pediatric extrapulmonary IMT is aneuploid. TP53 mutation and MDM2 amplification are rare (26). Because of the rarity of this proliferation in the terminal ileum, its natural history and biological potential are still uncertain (13), we reported a case of inflammatory myofibroblastic tumor in uncommon site and unusual clinical presentation with severe calcification and anemia.

IMT could not be distinguished clinically and radiologically from highly malignant neoplasm. The most important diagnostic aid is to bear this entity in mind when a child presents with intestinal obstruction associated with an abdominal mass. Radical surgical procedures or potentially harmful therapy should be avoided, and appropriate treatment is achieved by total excision of the lesion in most of the cases.
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Reference


