Recurrent Mesenteric Fibromatosis – A Case Report

Tejas A P1, Revanasiddappanadgir2, Harish Kumar3

1Junior resident, Department of general surgery, Raja Rajeswari Medical College and Hospital, Bangalore, Karanataka, India
2Professor, Department of general surgery, Raja Rajeswari Medical College and Hospital, Bangalore, Karanataka, India
3Assistant Professor, Department of general surgery, Raja Rajeswari Medical College and Hospital, Bangalore, Karanataka, India

Abstract: Mesenteric fibromatosis is a part of the clinical-pathologic spectrum of deep fibromatoses. The deep fibromatoses encompass a group of benign fibroproliferative processes that are locally aggressive and have the capacity to infiltrate or recur but not metastasize. Here we present a case of recurrent mesenteric fibromatosis in a 18 year old male patient. The Diagnosis is based on the histopathological examination of the tumor.

Keywords: Mesenteric Fibromatosis, Gardner’s Syndrome, Desmoid Tumor, Previous Abdominal Surgery, Enbloc Resection of the Tumor, Beta-Catenin Expression

1. Introduction

Mesenteric fibromatosis (MF) is a locally aggressive myofibroblastic proliferation of the mesentery. It is characterized by a spatially homogeneous proliferation of wavy spindle cells without atypia, associated with collagen deposition (often of the keloidal type), and an infiltrative border.

Mesenteric fibromatosis occurs in a wide age range of patients with 14–75 years of age. In contrast, abdominal fibromatosis occurs most commonly in young women, 20–30 years of age. 13% of patients with mesenteric fibromatosis have Gardner’s syndrome variant of Familial adenomatous polyposis.

2. Case Report

A 18 year old boy presented with complaints of mass abdomen and bilateral swelling of both the lower limbs since 2 years. Initially the swelling started as incidious in onset gradually progressive in nature to the present size, dull aching in nature, radiating to the back and lower limbs.

- Patient gives history of previous abdominal surgery for the similar complaints 3 years back.
- No history of fever/vomiting/altered bowel habits/burning micturion/ decreased urine output/jaundice.
- Patient had no comorbid conditions like diabetes, hyperstension, asthma.
- On examination, patient had pallor and bilateral swelling of the lower limbs.

On inspection of the abdomen there was a lower midline laparotomy scar of the previous surgery and a swelling was noticed in the abdomen located in the epigastric, left hypochondriac and hypogastric region measuring 15*10 cms not ballotable non pulsatile firm in consistency irregular surface and all borders are well made out and does not move with respiration and does not fall forward in knee chest position and dull to percuss and bowel sounds were heard. Hernialorifices and external genitalia were found to be normal. Digital rectal examination was found to be unremarkable. Other systems were normal.

Figure 1: Pre operative picture showing the mass in the abdomen and previous surgical scar

Patient was evaluated with working diagnosis of retroperitoneal tumour.

Patient was thoroughly investigated and all routine baseline investigations were found to be normal. Erect xray abdomen was normal.

CECT abdomen and pelvis was done as shown in the Fig 2 and Fig 3 shows the following changes:

Large lobulated moderately enhancing soft tissue density mass lesion measuring 22*16*12 cm in the abdomen and pelvis and retroperitoneal region encasing both mid ureters, sigmoid colon and few small bowel loops with bilateral moderate hydroureteronephrosis.

Colonoscopy was normal and Intravenous pyeolography was normal. With the above CECT showed features suggestive of desmoid tumor.

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Patient underwent exploratory laparotomy with intestinal resection and anastomosis under general anesthesia and Fig 4 showing the intra operative photograph of the tumor adherent to the sigmoid colon and small intestine and Fig 5 demonstrating resected tumor with part of the bowel weighting 4.2kgs.

Resected tumor was sent for histopathological examination and demonstrated as shown in Fig 6. Sprinkling of mononuclear cells and compressed thin walled vessels with plump endothelial cells and perivascular lymphocytic infiltrates. Features suggestive of desmoid fibromatosis.

Immunohistochemistry was done and spindle cells expressed beta-catenin as shown in Figure 7.
3. Discussion

Mesenteric fibromatosis is a type of fibroblastic proliferation affecting the mesentery that develops usually as a consequence of surgical trauma or occur spontaneously.

Imaging remains the mainstay of preoperative investigations to diagnose mesenteric fibromatosis.

The sonographic features of mesenteric fibromatosis are nonspecific and chiefly dependent on collagen and fibroblast content and intra-lesional vascularity of the tumor [1].

CT scan is considered the first-line imaging modality for identifying, characterizing, and staging fibromatosis.[2]

Microscopically, mesenteric fibromatosis is characterized by a spatially homogenous proliferation of wavy spindle cells without atypia, associated with collagen among dilated vessels. The mitotic count is relatively low with no evidence of necrosis and nuclear dedifferentiation [3].

MF does not express CD34 and S100 protein. Recently, the expression of beta-catenin was revealed in fibromatoses that might prove helpful.

Wide field surgical excision is the first-line treatment for most mesenteric fibromatosis [4]

Adjuvant radiation therapy reduces recurrence of mesenteric fibromatosis to 20%–40%, compared to 40%–70% with resection only [5].

Reference


