Large Retroperitoneal Desmoid-Type Fibromatosis of Renal Origin: Case Report and Literature Review

Tarun Jindal¹, Ankush Sarwal²*, Pravin Ramdas Pawar¹, Piyush Shankar Tripathi²

¹Department of Uro-oncology. Tata Medical Centre, Kolkata, India
²Department of Surgical oncology. Tata Medical Centre, Kolkata, India

*Corresponding author: Ankush Sarwal, Department of Surgical oncology. Tata Medical Centre, Kolkata, India. Tel: +91-9812636877; Email: drankushsarwal@yahoo.in


Received Date: 10 June, 2019; Accepted Date: 14 June, 2019; Published Date: 18 June, 2019

Abstract

Desmoid tumours are a rare type of tumours. Retroperitoneal desmoid is an uncommon mesenchymal neoplasm presenting as a firm mass with locally aggressive features. Desmoid tumour account for 0.03% of all tumours and 3% of all the soft tissue tumours. It usually manifests as a well-circumscribed or ill-defined, solid mass on cross-sectional imaging. The confirmatory diagnosis of these tumours is made by histological examination and immunohistochemistry. Surgical resection of the tumour is the treatment of choice in Desmoid tumours and also helps in clearing the diagnostic dilemma it creates in patient because of its rare nature. Here, we present a rare case of a Large Retroperitoneal Desmoid-type Fibromatosis of renal origin in a 28-year-old female, which was successfully treated surgically at specialised oncology centre in India.

Introduction

Desmoid-Type Fibromatosis (DF), also called desmoid tumour, is an uncommon mesenchymal neoplasm composed of fibrous soft-tissue proliferation [1]. The tumour is characterized by locally aggressive growth but rarely metastasize. Desmoid tumour may arise from anywhere in the body, but the common site being extra-abdominal (60%), abdominal wall (25%), and intra-abdominal (8-15%). Retroperitoneal DF is a rare clinical entity and accounts for less than 1% of retroperitoneal masses [2]. Desmoid tumour account for 0.03% of all tumours and 3% of all the soft tissue tumours. The incidence is 2.4-4.3 new cases per million people per annum. It occurs most often in the body between 20 to 35 years of age, with much more incidence in females as compared to males. Surgical resection of the tumour is the treatment of choice in DF and also helps in clearing the diagnostic dilemma it creates in patient because of its rare nature [3]. Here, we present a rare case of a Large Retroperitoneal Desmoid-type Fibromatosis of renal origin in a 28-year-old female, which was successfully treated surgically at specialised oncology centre in India.

Case Report

A 28-year-old female patient with a BMI of 21.35 kg/m², without significant surgical history or previous comorbidity presented to the outpatient clinic department with symptoms of nausea and vomiting for last 8 months, for which she has been taking symptomatic treatment form local practitioner for last 6 months. For last 2 months she has developed a lump in abdomen for which she had an ultrasonography of abdomen, which showed a 161 x 155 mm mass arising from the upper pole of left kidney. She could not recall any hospitalization or abdominal trauma in the past. On physical examination, there was hard lump palpable in the abdomen of size around 20 x 15 cm in the left lumbar region extending into Umbilical, Left hyochodrium and epigastric region. A CT scan (Figure 1) was done which showed a large lobulated heterogenous hypodense retroperitoneal lesion with enhancement within the lesion without calcification.
The lesion measured 21.9 x 18.9 x 10.3 cms. The lesion was not separate from the left kidney and suprarenal region and was displacing the spleen anteriorly and towards the left. Stomach was displaced to the right and pancreas was displaced anterior and surrounded by the lesion. Left ureter other retroperitoneal organs were normal without significant retroperitoneal lymphadenopathy. CT thorax which showed no thoracic metastasis. Ultrasound guided biopsy was done from the growth which favoured the growth to be desmoid tumour. Based on the symptoms as well as radiological and histological correlation, patient was planned for explorative laparotomy. Intraoperatively, there was a large retroperitoneal mass of size around 20 x 20 cm was seen occupying the whole left abdomen. Mass was seen infiltration the left kidney and spleen. The mass was densely adherent to distal pancreas and greater curvature of stomach. Based on these findings en-bloc removal of mass along with left nephrectomy, splenectomy, distal pancreatectomy and resection of wall of greater curvature was done (Figure 2). Primary repair of pancreas and stomach was done. One drain in sub hepatic space and one drain in pelvis was inserted and abdomen was closed in layers.

Histopathological examination of the specimen showed tumour of size 22 cm in greatest dimensions. Tumour is seen infiltrating the renal pelvis and abutting spleen, distal pancreas and gastric wall. All the margins were free of tumour. Histological sections showed a tumour composed of spindle cells, showing mild to moderate atypia, arranged in fascicles with interspersed small arterioles. Mitotic count less than 1/10 high power field. Immunohistochemistry showed tumour cells to be positive for B-catenin, SMA and desmin, while negative for S100 and DOG1, suggesting it to be desmoid-type fibromatosis.

Postoperatively, patient was shifted to intensive care unit for post-operative management for 3 days and kept nil by mouth for 3 days with intravenous parental nutrition and albumin. Consequently, the patient was gradually shifted to oral liquid diet then soft diet. Sub hepatic drain and pelvis output was reported initially high and was clear in colour. Gradually, the drain output reduced in amount to less than 50 cc per day. Sub hepatic drain was removed on 5th post-operative day and pelvis drain was removed on 7th post-operative day. The patient recovered well and was discharged on the 8th post-operative day. Post-operatively in view of R0 resection and no distant metastasis, no further adjuvant therapy was given to the patient. The patient has been asymptomatic for 12 months, on clinical and radiological evaluation she is recurrence free and living a healthy normal lifestyle.

Discussion

Desmoid-Type Fibromatosis (DF) also called desmoid tumour or musculoaponeurotic fibromatosis, is an uncommon mesenchymal neoplasm composed of fibrous soft-tissue proliferation with the production of intracellular collagen [1]. The disease was first described by Mueller in 1838 as a band, or a cyclic structure, and the term fibromatosis as a group of similar changes, was first used by Stout in 1961 [1, 4]. Desmoid tumour may arise from anywhere in the body, but the common site being extra-abdominal (60%), abdominal wall (25%), and intra-abdominal (8-15%). Retroperitoneal DF is a rare clinical entity and accounts for less than 1% of retroperitoneal masses [2]. Desmoid tumour account for 0.03% of all tumours in literature and 3% of all the soft tissue tumours. The incidence is 2.4-4.3 new cases per million people per annum. The incidence of DF is common in young females between 3rd and 4th decade of life. DF has been associated with multiple risk factors including previous surgery, genetic mutations of the adenomatous polyposis coli gene such as in FAP or the beta-catenin gene, trauma, pregnancy, and oral contraceptive use. However, the pathogenesis of DF is not completely understood [3]. The patient can be asymptomatic or could present with vague abdominal symptoms and abdominal lump. On Radiological imaging DF appears as well-defined solid mass of variable attenuation and enhancement, which depends on tissue components. DF with a highly collagenous stroma usually displays homogeneous, soft-tissue attenuation on CT scans. The soft-tissue component can often mimic similar to other solid tumours, such as gastrointestinal stromal tumor, lymphoma, or...
soft tissue sarcoma [5]. Imaging exams are useful in establishing size, extension and anatomical relationship, the confirmatory diagnosis of these tumours is made by histological examination and immunohistochemistry. The cells usually have a poorly circumscribed pattern with spindle cell proliferation forming long beams or spiral patterns and are strongly positive to vimentin staining and the immunoreactivity to beta-catenin is expressed in the 67-80% of cases. Immunohistochemically, the tumour cells are negative for CD34, CD117, and S-100 protein [6] The main therapeutic strategy is to prevent the invasion of surrounding tissues. The therapeutic decision requires the approach of a multidisciplinary team. Surgery is considered the treatment of choice whenever possible, other alternatives include radiotherapy, hormone therapy, treatment with NSAIDs and even observation have also been reported. In a study done by cates et al, a positive or close (<1 mm) resection margin was an independent prognostic factor for local recurrence in a multivariate Cox regression model accounting for patient age and anatomic site (hazard ratio 9.52; P=0.028), thus emphasizing the need for R0 resection in such cases [7]. In present case Desmoid-type Fibromatosis had renal origin and which has been rarely reported in literature.

**Conclusion**

Desmoid tumours are rare clinical entity. We report a rare case of a Large Retroperitoneal Desmoid-type Fibromatosis of renal origin in a 28-year-old female, which was successfully treated surgically at specialised oncology centre in India. Surgical resection of the tumour is the treatment of choice in DF and also helps in clearing the diagnostic dilemma it creates in patient because of its rare nature and requires a multidisciplinary team approach.

**References**