An aggressive fibromatosis in the palm of a female child: A rare case

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ABSTRACT

Aggressive fibromatosis is the locally aggressive benign tumor of mesenchymal origin. It can be found in any part of the body. However, abdomen is the most common site of this lesion. It has a predilection to females between 15 and 60 years. We report the case of a 16-month-old female child with an aggressive fibromatosis in her right palm. The location and age of presentation make this a rare case. She was diagnosed by histopathological examination and the mass was excised. The patient was advised for follow-up examination due to the high degree of recurrence of this tumor.

Key words: Aggressive fibromatosis, Collagenous stroma, Trucut biopsy

Computed tomography (CT) scan of the right palm was ordered that showed a multilobulated hypodense soft-tissue lesion of approximately 22 × 46 × 54 mm size along the palmar aspect of the right hand (Fig. 2a and b). There was encasement of muscles and tendons along the palmar aspect with loss of fat planes, representing probably as a fibrosing tumor.

A trucut biopsy of the lesion on the right palm was done and was sectioned and stained with hematoxylin and eosin stain. The microscopy showed hypocellular fibrous tissue revealing spindle or stellate cells widely separated by abundant collagenous stroma. Few adnexal structures and fragmented epidermis were also noted (Fig. 3a and b). There was no mitosis, nuclear atypia, inclusion, hyaline deposit, or vasculitis favoring the histological diagnosis of fibromatosis (Fig. 3c and d).

The patient was admitted to daycare. After admission, general examinations including vitals and necessary blood investigations (Comprehensive Metabolic Panel, Human Immunodeficiency Virus, Hepatitis B surface antigen, and Hepatitis C virus) were done and found normal. She underwent the excision of the mass under general anesthesia. Volar approach (modified Henry extension) was done and the lesion was marginally excised. Ulnar nerve and median nerves were protected and were prophylactically decompressed in the Guyon canal. Hypothenar muscles (abductor digiti minimi, flexor digiti minimi, and opponens digiti mini) were completely encased in mass, so they were removed with the tumor. The remaining thenar muscles, flexor digitorum superficialis, and flexor digitorum profundus were protected.

Postoperatively, she was shifted to the daycare and managed with cefuroxime 250 mg BD, ibuprofen 5 ml BD, pantoprazole 20 mg OD, intravenous fluids, and other supportive measures.

CASE REPORT

A 16-month-old female child presented to the outpatient department with complaints of the right palm swelling for 1 year. The swelling was insidious in onset that gradually increased in size, ultimately covering the whole palm area. The patient had no complaints of fever, pain, and trauma. There were no significant past and family histories.

On arrival, the patient vitals were normal. On examination, there was a firm swelling covering the entire palmar surface of the right hand (Fig. 1a and b). No local inflammatory signs were found.

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Nil by mouth was cleared and the patient tolerated the oral diet well. She mobilized well, with no fresh complaints. Hence, given her stable vitals, she was discharged after explaining further care and follow-up advice. Although she was advised to come for follow-up at 1 month, 3 months, 6 months, and follow-ups have to be maintained for at least 2 years for recurrence; the patient came for follow-up visits for only 3 months. She was able to do her activity of daily living. She had complete atrophy of hypothenar eminence but was able to do remaining hand function.

DISCUSSION

Aggressive fibromatosis, also called a desmoid tumor, is a locally aggressive tumor that originates from fibroblast cells. These cells are a part of the connective tissue. Since connective tissues are present everywhere in the body, aggressive fibromatosis can occur in any part of the body. However, the abdomen is the most common site of presentation [5]. Aggressive fibromatosis is not encapsulated with a poorly defined margin. Hence, it has a high tendency to invade local tissue and recur after treatment [6]. Local invasion to the neurovascular structure can cause pain and loss of function [4].

Although the exact etiopathogenesis of aggressive fibromatosis is still unknown, it is considered to be sporadic or hereditary. Trauma and surgical scar sites have a higher tendency to develop this condition. Similarly, pregnancy and oral contraceptive use is also associated with the development of aggressive fibromatosis [4]. Furthermore, studies have shown that this condition has more predilections toward females [5]. Association of the desmoid tumor with hereditary conditions such as Gardner syndrome and familial adenomatous polyposis (FAP) supports the hereditary nature of this disease [3]. Approximately 85% of sporadic cases show a mutation in the beta-catenin pathway. It is known that 45F mutation in the beta-catenin pathway is associated with an increased lesion recurrence rate. Similarly, desmoid tumor associated with FAP shows a mutation in the APC gene. Moreover, this desmoid tumor has a higher tendency to grow at the surgical scar site [6].

Ultrasound, CT scan, and magnetic resonance imaging (MRI) are used for the imaging of aggressive fibromatosis for investigation and follow-up. They help define the origin and extent of local invasion of the lesion and guide the surgery plan [6]. Although MRI is more sensitive, CT scan is commonly used for imaging this tumor [4]. It presents as soft-tissue mass with poor margin and infiltration of the surrounding tissue. Metastasis is always absent here [3]. The diagnosis of aggressive fibromatosis is made by histological findings. It presents as the proliferation of spindle-shaped fibroblast cells, massive collagenous stroma, and blood vessels without nuclear atypia and hyperchromasia. It also stains positive for beta-catenin, vimentin, cyclooxygenase-2, tyrosine kinase, and estrogen receptor beta [6].

Surgery is the mainstream treatment for aggressive fibromatosis. However, it is associated with recurrence and a high degree of morbidity and mortality [7]. The limitation of surgery can be overcome by radiotherapy alone or radiotherapy along with surgery as a treatment modality [8]. Other treatment options for aggressive fibromatosis are chemotherapy (methotrexate, tyrosine kinase inhibitor, and vinblastine), hormonal therapy (tamoxifen and raloxifene), non-steroidal anti-inflammatory drugs, high-intensity focused ultrasound treatment, and ablation technique (radiofrequency ablation) [1].

Recurrence is common in aggressive fibromatosis. Although age, size, and site of the tumor have no association with recurrence, a positive margin has a high chance of recurrence. Hence, a follow-up for at least 2 years with a wait and watch policy followed by adjuvant radiotherapy is offered in case of recurrence [9].
CONCLUSION

Aggressive fibromatosis in the palm of a 16-month female child is a rare case in terms of the lesion site and age of presentation. Due to the lack of capsule and its nature of the aggressive local invasion, the patient has to be followed up for the possibility of recurrence.

REFERENCES


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