Pediatric Nodular Fasciitis: Case Series Report

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Abstract

We like to report on two paediatric cases presenting with Nodular Fasciitis. Both were rapid growing soft tissue masses excised between Jan 2016 to Dec 2016. After surgical excision they were followed up in outpatient, we would like to describe the histological evaluation and management of these cases.

Keywords: Nodular Fasciitis; Inflammation; Sarcoma

Introduction

Nodular Fasciitis (NF) is a rare benign soft tissue tumor more commonly seen in the adult, compared to the pediatric population. It is often mistaken as a malignant soft tissue sarcoma. Benign histology changes the course of management strategy to less aggressive therapy. Our institution reports two pediatric cases of NF, presenting with a soft tissue mass clinically thought to be a soft tissue sarcoma.

Case Series

Case presentation 1

One year old girl presented, with 3 month history of fast growing mass on the forehead. Two days prior to the onset of growth there was history of blunt trauma to the forehead. The child was born at term, and past medical history was unremarkable. Clinically the mass had a smooth surface, mobile and not fixed to the underlying skull. No audible bruit was present. Neurological examination was normal. The ultrasound showed a complex predominantly solid mass with cystic components and micro-calcifications. Doppler ultrasound revealed moderate vascularity with arterial waveform in the solid component. MRI scan showed right lateral supraorbital mass measuring 39 mm x 46 mm x 23 mm. The mass was superficial to the skull, preseptal and not involving the orbital structures.

Surgical excision was performed. A wide coronal incision was made 2 cm behind the hairline, dissected to the subgaleal fascia plain avoiding the temporal branch of the facial nerve (Figure 1), and supraorbital nerve. The lesion was adherent to the frontalis muscle posteriorly and temporo-parietal fascia. There was no definitive capsule and therefore the lesion was excised in pieces rather than intact. The specimen was submitted for urgent histological evaluation. No complications occurred post-operatively (Figure 2).

Case presentation 2

10 year old girl of mixed race, presented with a one month history of a progressive enlarging non-painful mass on the left anterior chest wall. There was no preceding history of trauma prior to presentation. Anatomically the mass was located at the fourth intercostal space, mid clavicular line. It was mobile, not adherent to any surrounding structures, no audible bruit was detected. Chest x-ray was normal. The mass was surgically excised one week after presentation. No imaging was done prior because of its small size. The patients were followed up at 2 weeks and 3 months post operatively, no complications or recurrence occurred.

Histology

Microscopic sections show a tumor comprising plump myofibroblastic cells growing in a vaguely storiform pattern. These cells had vesicular nuclei with small nucleoli and basophilic to eosinophilic cytoplasm. Mitotic activity was present. Randomly arranged within the tumor were collagen fibers and osteoclast-like giant cells (Figure 3). Myxoid foci were prominent in some areas. The lesion was...
seen infiltrating peripheral soft tissues. Immunohistochemical stains showed positive staining for smooth muscle actin, which appear very different to a malignant lesion (Figure 4). Desmin, CD10, CD34 and S100 were negative. NF was extending to the excision margin.

Discussion

Nodular fasciitis is a rare benign lesion, consisting of proliferating fibroblasts. It is usually in the deep fascia in the head and neck areas, its rapid growth can be deceptively similar to that of soft-tissue sarcomas [1,4,5]. Approximately 10% of all known cases are found in children [1,4]. Male to female ratio is equal. The lesion is an idiopathic self-limiting reactive process of unknown etiology. The lesion is described as being a trauma-related phenomenon [1,4,5]. Differential diagnosis consists of solid malignancies: rhabdomyosarcoma, neuro blastoma, osteosarcoma, neurofibroma, lymphoma and cysts/abscess [2] (Figure 4). Clinically it presents as a progressively enlarging painless solid mass. The clinical symptoms are non-specific and do not help with the pre-operative diagnosis [3]. Impingement of the surrounding structures can produce pain or other symptoms. Constitutional symptoms such as fever and weight loss are rare [4]. Microscopic evaluation shows discrete and nodular hyper cellular growth of fibroblasts and myofibro blasts centered in the tissue. Prominent mitotic activity, areas rich in ground substance with amyloid and microcystic change, and extravasates erythrocytes and inflammatory cells are often present. These characteristics are the classical description of NF [1].

Radiological imaging includes sonar, CT or MRI [1]. Ultrasound helps in differentiating solid tumors from cystic lesions, specifically vascular malformations and post-traumatic false aneurysms. MRI is the imaging modality of choice because it helps to define the lesion in relation to surrounding vital structures, to map surgical approach and detect associated bone remodeling or destruction [1]. The signal characteristics are variable depending on the histological components.

Cellular lesions show T1 Hyperintensity and T2 Hyperintensity due to the myxoid component, whereas more collagenous tumors are hypointense on both sequences. The mass typically enhances diffusely on T1 post-contrast sequences [7,8]. Complete surgical excision is the definitive treatment. The lesion is usually unencapsulated. Observation or partial resection may be considered in special cases [1]. The condition is a self-limiting process with a low recurrence rate [1]. The effectiveness of use of steroids in NF is not known. Cranial fasciitis is a histologically similar condition to nodular fasciitis with involvement of the outer table of the skull bone. One report advocates local intra-lesional injection in this subtype of NF should it recur [6].

References