Superficial acral fibromyxoma. Un uncommon tumor of the foot
Fibromixoma acral superficial. Um tumor incomum do pé

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ABSTRACT
This study presents a case report of a 52-year-old female patient with a superficial acral fibromyxoma (SAF) of her left foot fifth toe. This is a very uncommon soft tissue tumor that affects mainly hands and feet, with a special predilection for toes, with the hallux most frequent. The tumor in this case was 3cm long and 2.5cm wide, which was unusually large. Few cases are larger than 2cm long and localized on the fifth toe. The treatment was wide resection of the tumor through amputation of the fifth toe.

Level of Evidence V; Therapeutic Studies; Expert Opinion.

Keywords: Fibroma/surgery; Soft tissue neoplasms; Toes/surgery.

INTRODUCTION
The superficial acral fibromyxoma (SAF) is a very uncommon soft tissue benign tumor. It is also known as digital fibromyxoma or cell digital fibroma. It can occur at any age, typically involving fingers and toes, with particular predilection for the hallux. It has a very slow growing pattern, with imaging studies showing soft tissue compromise with or without bone erosion. Given its recurrence potential and its location, treatment is surgical resection with wide margins.
**CASE REPORT**

This study was approved by the Research Ethics Committee of our institution as well as the informed consent signed by the patient.

The patient was a 52-year-old obese female, without any other relevant history, who complained of a mass at the distal aspect of her left fifth toe. There was a two-year history, with a slow growing progression and slight pain. She related mild previous trauma to her foot 2 years before. On physical exam, she had a lobulated mass of approximately 3.5cm diameter, which eliminated and deformed the nail region. The surface was wrinkled with firm consistency and defined limits (Figure 1 A and B). The remainder of the physical examination was normal. Foot radiographs showed a soft tissue mass with slight bone erosion or scalloping at the distal phalanx region (Figure 1 C). There were no signs of malignancy. Chest radiographs were normal. The MRI showed a 30mm diameter mass on the dorsal aspect of the fifth toe, with lobulated borders. It showed medium intensity signal in T2 and T1, and a heterogeneous appearance with some hyperintense regions. The mass could be seen in contact with the cortex of the distal phalanx, which showed an irregular border (Figure 1 D-F). Amputation of the fifth toe was performed with wide margin. The tumor was sent for pathological analysis (Figure 1 G-I).

On the macroscopic analysis, a 30mm diameter mass, with dermis and subcutaneous tissue compromise was observed. On microscopic analysis, it featured benign mesenchymal cell proliferation with a multinodular morphology of expansive growth. Each of these foci shows a myxoid matrix with thin collagen fibers, stellate cells without atypia, and slight vascularization. A good surgical margin was obtained from the entire tissue resected (Figure 2 A and B). The immunohistochemistry (IHC) showed markedly positive reaction for CD34 and CD10 and negative for EMA. It also showed negative reaction for

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**Figure 1.** Clinical and radiological illustrations. A) Dorsal view. B) Plantar view. A 3.5cm lobulated tumor that eliminates and deforms the nail region can be observed. C) Dorsal view x-ray. A soft tissue mass can be observed distal to the distal phalanx of the fifth toe. D) MRI: T1 sequence. Sagittal plane. E) MRI: T2 sequence. Sagittal plane. F) MRI: T2 sequence. Coronal plane. Medium signal is observed in every sequence with some hyperintense regions. G and H) Trans-proximal phalanx amputation of the fifth toe dorsal and lateral view. I) Resected tumor.
S100 Protein, Chromogranin, and Actin which ruled out Schwan cell tumor, glomus tumor, and a tumor of smooth muscle origin respectively (Figure 2 C and D). The clinical findings, location, imaging studies, morphology and IHC profile, confirmed a superficial acral fibromyxoma. At 14 months of follow-up, she has no pain, no signs of relapse and good functional outcome.

DISCUSSION

SAF was first described by Fetsch et al.\cite{3}, in a retrospective study of 37 cases. It affects males and females with male predominance 2:1. It can occur at any age (range, 4-86 years), with an average age of 48. It typically compromises the nail bed (subungual region) in more than 95% of cases according to Hollomann\cite{2}. Less commonly, the tumor has been found in the palm and sole, the heel, or more rarely in the leg and thigh. The most frequent involvement site is the hallux\cite{3,4}. SAF typically shows a chronic behavior, characterized by a slow growing mass with little or no pain at the distal aspect of a finger or toe. Nail deformation is common because of the nail bed compromise. Given the slow progression and the limited symptoms of this tumor, medical consultation is delayed in the majority of cases. In Fetsch’s retrospective series, time from appearance to first medical consultation varied from 3 months to 30 years with an average of 3 years. Our case presented with a two years history. SAF has been described with varying shapes including unilocular and multilobulate. It seldom compromises mobility, and has little pain to palpation\cite{3}.

On routine radiographs, SAF appears as a soft tissue mass with or without bone compromise. This could be scalloping or sclerosis of the underlying bone, which is present in more than 30% of cases\cite{2}. In contrast, our case showed a slight bone erosion at the distal aspect of the phalanx. On MRI, a hyperintense image could be observed on the T2 images. SAF typically appears as a solitary lesion growing mainly in dermal or subcutaneous tissue areas and less commonly reaching the deep fascia layer.

Histologically, it shows a spindle cell and stellate proliferation, enclosed in a myxoid or collagen matrix. Some regions may show a storiform pattern and blood vessels and stromal mast cells are often observed. Mitotic figures are usually sparse, averaging <1 per 10 HPF\cite{5}. SAF shows multinucleate cells in 50% of cases but with minimal nuclear atypia. Immunohistochemistry typically shows positive reaction to CD 34 in every case, while epithelial membrane antigen (EMA) and CD99 are positive in many cases. S100 protein, glial fibrillary acid protein (GFAP), actin, desmin and keratin are typically negative in the SAF\cite{2,3}.

Given its rare appearance, SAF is often mistaken for other myxoid soft tissue tumors. The differential diagnoses can be classified into benign or malignant tumors. In the benign myxoid group, periungual fibroma and acquired digital fibrokeratoma are among the most common differential diagnoses. They typically extend into the dermis and have more cellularity on microscopy\cite{6,7}. Neurofibroma can also be part of the differential diagnosis although it shows an S100 Protein expression. In the malignant myxoid group, dermatofibrosarcoma protuberans (DFSP), low-grade fibromyxoid sarcoma (LGFMS) and myxofibrosarcoma are the main differential diagnoses\cite{8}. DFSP predominately presents on the trunk, with an extremely low incidence localized to the hands and feet, and it is negative for EMA\cite{2}. The LGFMS has a more aggressive behavior and disorganized cell growth pattern than SAF. It affects deeper structures and acral localization is uncommon\cite{8}.

In his series of 37 patients with SAF, Fetsch et al. described two cases with significantly more focal atypia and a low mitotic rate, which suggested that these cases might become low-grade malignant neoplasms, although there was no evidence of an aggressive progression of these two tumors\cite{3}. In order to reduce the risk of recurrence, which is low, a wide margin at the resection site is recommended\cite{3}. In the study by Holloman et al., the recurrence rate was 24% with an average follow-up of 27 months. However,
there were no recurrences in those cases with a free margin at the resection\(^2\).

We have a follow-up of this patient of 14 months with excellent clinical results. The clinical history and examination findings in this patient, together with the clear imaging studies, histology and IHC, provide a unique study of this rare soft tissue tumor. Localized to the fifth toe, and 3cm long x 2cm wide, this case was quite unusual in light of the less than 170 cases previously published in the literature\(^9\).

**CONCLUSION**

We have shown a case of a superficial acral fibromyxoma of the fifth toe of the left foot of a 52-year-old female patient. While this is an uncommon, benign, slow growing, soft tissue tumor, it has a predilection for fingers and toes as well as hands and feet. It should be part of the differential diagnosis of foot and toe tumors of myxoid origin. Surgery is the treatment of choice and normally curative with wide resection. There are no documented instances of metastases.

**REFERENCES**