Calcifying aponeurotic fibroma: a case report
Fibroma aponeurótico ossificante: relato de caso

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ABSTRACT
Calcifying aponeurotic fibroma (CAF) is a rare lesion that can affect the feet and should be considered as a differential diagnosis of plantar tumors. The treatment is given by surgical resection, and the definitive diagnosis is confirmed by histopathological analysis. This case report shows a 17-year-old patient with CAF, the concept of the treatment performed, and emphasizes the importance of proper imaging planning to reduce risks and recurrence.

Level of Evidence V; Therapeutic Studies; Expert Opinion.

Keywords: Fibroma; Foot; Tumor; Surgery.

INTRODUCTION
Calcifying aponeurotic fibroma (CAF) is a rare benign tumor of fibroblastic origin with locally aggressive behavior(1) that typically develops in the fascia and tendons of the palms or soles of children and adolescents. Although it has been described in several age groups, the peak incidence occurs between 8 and 14 years of age and it is twice as frequent in males as in females(2-5). The main sites of involvement are the hands (77% on palms and fingers) and the feet (13%)(6). The typical presentation of the tumor is as a painless, locally aggressive, subcutaneous, poorly defined, firm mass of slow growth that is not adherent to the skin, but to the adjacent subcutaneous tissue(6). Malignant transformation is rare, though change into fibrosarcoma has already been described(2,7), as well as metastases to bone and lungs. The lesion is not easily identified on radiographs. The most recommended supplementary assessment for diagnosis and preopera-
tive planning is magnetic resonance imaging (MRI), as it enables evaluating the extent of lesion margins and the complexity of calcifications.\textsuperscript{[1,7,8]} Differential diagnoses of CAF include synovial sarcoma, undifferentiated pleomorphic sarcoma, mainly in the hands and wrists\textsuperscript{[9]}, epithelioid sarcoma, clear cell sarcoma, giant cell tumor of tendon sheath, soft tissue chordoma, palmar or plantar fibromatosis, Morton's neuroma, and peripheral nerve sheath tumor (Schwannoma). The recommended treatment is complete surgical excision including the oncological margins\textsuperscript{[6,7]}, followed by a histological analysis and an immunohistochemical analysis\textsuperscript{[8]}. Due to the locally invasive pattern, after surgical excision, CAF has a high rate of recurrence.\textsuperscript{[1,3,5]}

The purpose of this study is to present a case report of CAF and the concept of the treatment performed, as well to present a literature review on CAF to highlight the possibility of this specific diagnosis considering the differential diagnoses of tumors that affect the foot.

**CASE REPORT**

This study was approved by the Ethics Committee, being registered on Plataforma Brasil under CAAE number 23237719.7.0000.5404, and follows Brazilian and international guidelines on ethics in research involving human participants. An informed consent form was signed.

The patient was a 17-year-old male adolescent complaining of sporadic pain and tumor in the plantar region of the left forefoot for seven years. There was no history of trauma and the condition was slowly progressive, with pain worsening with physical activity. Upon examination, a tumor of moderately hard consistency was found in the plantar region, below the first metatarsal and appose to the lateral sesamoid. Interphalangeal hallux valgus was present, with no skin changes, neurovascular changes, or instability of the first ray. The initial clinical suspicion was a relatively deep and adherent tumor or fixed sesamoid dislocation. Radiographs and MRI of the left forefoot were requested.

On the initial radiograph (Figure 1), it was possible to observe an enlargement of the space between the first and second metatarsals of the left foot, normal positioning of the sesamoids, and interphalangeal hallux valgus. It was further possible to notice small proximal calcifications between the first and second metatarsals.

On MRI (Figure 2), an elongated, lobulated tumor lesion was identified adjacent to the plantar contour of the lateral sesamoid of the hallux, with multiple intratissenal calcified foci. The tumor extended proximally to the medial belly of the flexor hallucis brevis and underlying plantar fat, measuring 4.0cm long x 1.9 cm deep x 1.8cm wide. The signal was predominantly low on T1 and T2, with evidence of enhancement after the administration of paramagnetic contrast, especially in late sequences. There was also another small lesion with the same characteristics next to the plantar and lateral contour of the first tarsometatarsal joint, next to the insertion of the peroneus longus tendon. Potentially, the tumoral lesion showed no signs of aggressiveness, which is compatible with the time of clinical evolution, and a possible vascular origin with calcified/fibrous component was considered as a diagnostic hypothesis.

The treatment performed was the complete surgical excision of the plantar mass in the forefoot, adjacent to the lateral sesamoid, including resection of the sesamoid. During resection, the tumor extension to the flexor hallucis longus was confirmed. The tumor affected the tendon and the fibroadipose and muscle tissues, extending to the periosteum of the first metatarsal without, however, infiltrating it, thus the margin of this bone was not to resected. The anatomopathological analysis identified a fasciculated, sclerosing, infiltrative, moderately cellular spindle cell proliferation, with the formation of calcified nodules of chondroid matrix, calcified metaplasia, and cartilaginous and bone metaplasia, defining the diagnosis as CAF.

At the initial follow-up, the patient remained without symptoms, pain, or new tumors, resuming the practice of...
physical activities after wound healing. On control radiograph four months after surgery, an increase in the proximal space between the first and second metatarsals persisted. On control MRI (Figure 3) six months after surgery, there was a change in the plantar and lateral signal to the first metatarsal at the site of the surgical scar, but without clear evidence of recurrence.

One year after surgical resection, the patient remained asymptomatic, however, on the control MRI (Figure 4), tumor recurrence was observed with characteristics similar to the first examination. The lesion was measuring 1.1 x 0.9 x 0.6cm. Besides, changes to the tissue that infiltrated the first intermetatarsal space were more evident and involved the Lisfranc ligament, with larger components in the plantar and lateral portions.

As the patient remained asymptomatic, with no changes in daily activities, without restriction of movements or other complaints, and without palpable masses, it was decided to perform control MRIs and maintain clinical follow-up, which to date totals 19 months.

**DISCUSSION**

Calcifying aponeurotic fibroma is a rare benign tumor that affects mainly the hands and feet of children and adolescents. It has already been described in other body parts, such as the mandible, neck, scalp, forearm, thigh, knees, abdomen, and lumbosacral region\(^7\). Keasby first described it in 1953 as a juvenile aponeurotic fibroma, also called juvenile fibromatosis\(^7\). Patients present poorly defined, hard, small masses of soft tissues of slow growth\(^5\). The epidemiological presentation of our case is quite typical: a 17-year-old male adolescent with plantar tumor of slowly progressive growth since he was ten years old.

On radiographs, it is possible to find multiple or speckled calcifications and bone erosions\(^4\), although cortical erosion is rarely seen\(^9\). At the initial radiographic examination, there were thin plantar calcifications in the first intermetatarsal space up to the Lisfranc joint and an enlargement between the first and second metatarsals. However, such radiograph is not enough for a definitive diagnosis. MRI and computed tomography are the preferred examinations to diagnose and define the lesion. Tomography may reveal an unspecific soft tissue mass with dotted calcification and may show the infiltrative growth pattern of the lesion into surrounding tissues\(^4\). On MRI, the findings include poorly defined soft tissue mass with speckled calcifications with a low or intermediate intensity signal on T1 and an image of intermediate to high intensity on T2. On T2, areas of heterogeneity are also observed, with few areas of low or intermediate intensity\(^4\), being affected by the degree of calcification, reduced cellularity, and quantity of fibrous components\(^4\). As demonstrated, the homogeneous image shows an increased signal after injection of gadolinium contrast\(^9\), which may be diffuse or with peripheral enhancement\(^9\).
In the present case, as there was no evidence of malignancy in the images and considering the history of slow and progressive growth, a surgical excision of the tumor was performed. The lesion extended to the medial belly of the flexor hallucis brevis, with multiple foci of calcifications inside of it. Although it is located in the subcutaneous layer in about 80% of cases, CAF can present inter or intramuscular involvement\(^4\). The exact limits were not evident during our surgical excision, and as the tumor involved the lateral sesamoid, it was decided to resect it en bloc with the affected soft parts. However, during surgery, it was not identified whether the lesion extended proximally to the first tarsometatarsal joint or if there was an isolated focus other than the proximal lesion. No frozen-section biopsy of the margins was performed during surgery due to the benign nature of the lesion.

The definitive diagnosis is based on histological findings, immunohistochemical tests, and ultrastructural...
studies. On histopathological analysis, CAF has a biphasic morphology, with moderate and infiltrative cellularity, a component similar to fibromatosis, and calcification nodules accompanied by rounded epithelioid cells, with infrequent mitotic figures. The present case presented similar findings, with moderately cellular, infiltrative spindle cell proliferation, formation of nodules of chondroid matrix, and cartilaginous and bone metaplasia.

The infiltrative nature of CAF is manifested by high rates of recurrence, higher than 50%, which can take years after the initial excision\(^1\)\(^2\)\(^5\). The risk of local recurrence appears to be higher in younger children, and, in general, recurrence is not destructive\(^1\)^\(^3\)^\(^5\). In this case report, a tumor was observed about a year after surgery in a patient in late adolescence. It was not possible to conclude if there was a tumor remaining due to incomplete surgical resection or if it was an earlier recurrence of the CAF comparing with what is described in the literature.

**CONCLUSION**

In conclusion, CAF is a rare lesion that can affect the feet and should be considered as a differential diagnosis of plantar tumors. It is treated by surgical resection and its definitive diagnosis is confirmed by histopathological analysis, there being high rates of recurrence. The present case highlights the importance of a proper planning with images to reduce risks and postpone recurrences.

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**Authors’ contributions:** Each author contributed individually and significantly to the development of this article: LBO *(https://orcid.org/0000-0002-8606-6187) conceived and planned the activities that led to the study, wrote the article, interpreted the results of the study, participated in the review process, approved the final version; DM *(https://orcid.org/0000-0002-3893-0292) participated in the review process, approved the final version; KVG *(https://orcid.org/0000-0001-5025-0823) interpreted the results of the study, participated in the review process, approved the final version; MCMD *(https://orcid.org/0000-0001-6572-1771) conceived and planned the activities that led to the study, wrote the article, interpreted the results of the study, participated in the review process, approved the final version. *ORCID (Open Researcher and Contributor ID).*

**REFERENCES**