Leiomyoma of the calf muscle in a child with calcification and ossification – a case report

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Abstract
Leiomyomas (smooth muscle tumours) arising in skeletal muscle are extremely rare in children, especially in the extremities.
A 3-year-old girl presented with a slow-growing mass in the left calf which had been present since the age of 6 months. Plain radiographs revealed a well-circumscribed mass, fusiform in shape, not attached to bone. There was widespread calcification of the tumour. MRI confirmed extensive calcification and ossification in the soleus and medial head of the gastrocnemius displacing the neurovascular bundle anteriorly. A diagnosis of a sarcoma was suggested.
The mass was well defined at surgical exploration, encapsulated and easily dissected from the soleus, gastrocnemius muscles and blood vessels.
The tumour was gritty on cutting with a saw and histology confirmed a leiomyoma of the soleus muscle. Immunohistochemical markers for smooth muscle actin and desmin were positive. The recovery was uneventful at five-month follow-up.

Key words: leiomyoma, calf muscle, calcification, ossification, immunohistochemistry

Introduction
Leiomyomas are extremely rare in the deep soft somatic tissues of the body. Those described have been seen mainly in the cutaneous and subcutaneous tissues and in the genito-urinary tract and abdominal cavity mainly in adult females. Leiomyoma of deep soft tissue, arising in skeletal muscle of the extremity, is rarely seen by an orthopaedic surgeon. The lesion may be calcified or even rarely show ossification with calcification. These radiological features can be misdiagnosed for more familiar conditions like calcified tuberculous nodes, calcified haematoma, myositis ossificans, tumoral calcinosis or benign and malignant tumours of soft tissue seen in the musculoskeletal system. Normal smooth muscle is widely distributed in the body being present in the walls of the genito-urinary, gastrointestinal, respiratory tracts, skin and blood vessels. Leiomyoma arising in skeletal muscle is thought to be of vascular origin. Yannopoulos and Stout expressed a probable venous origin. Goodman suggested that these tumours arose from undifferentiated mesenchymal cells or smooth cell rests. Kransdorf in a ten-year review of benign soft tissue tumours in about 40 000 cases found no deep leiomyomas in children <14 years old.
Farman in a review of 7,748 leiomyomas found that 95% occurred in the female genital tract and the remainder were distributed over various sites. Billings in a large study found only three children with deep soft tissue leiomyomas, two involved the forearm and one in the thigh. Eleven cases involving the extremities in children have been described in the literature and seven cases involving skeletal muscle have been reported. Calcified lesions have been described on six occasions. Calf muscle involvement has been reported in two children previously. Ossification with calcification was seen in one case only.

The aim of this report is to increase the awareness of the clinicopathological features of leiomyoma of deep soft tissue in children, and to include the tumour in the differential diagnosis of diffuse calcification in soft tissue in children.

Case report
A 3-year-old girl presented with a swelling over the posteromedial aspect of the left calf. The swelling was noticed at 6 months and gradually increased in size. Pain and a limp were the initial presenting features. There was no history of trauma. On clinical examination there was a fusiform swelling over the posteromedial aspect of the left leg in the calf muscle ± 10 cm × 6 cm (Figure 1).

The surface temperature was normal, the mass was smooth, firm with well-defined edges, not attached to bone or skin, with no fluctuance. Pain was elicited on pressure, and the child walked with a limp. The tendo-Achilles was contracted and dorsiflexion to neutral was elicited. There was no neurovascular deficit. No local or generalised lymph nodes were palpated. The child was apyrexial. The plain radiographs of the left leg showed an oval, well-circumscribed lesion in the calf with calcific densities and areas of opacification (Figure 2). A rim of calcification around the lesion suggested a pseudo capsule. Magnetic resonance imaging (MRI) with intravenous contrast showed an oval mass in the middle and distal calf, with no bony involvement, in the posterior compartment (Figures 3a and b). The mass had a heterogeneous signal intensity and demonstrated intense enhancement. Areas of ossification were demonstrated. The lesion demonstrated expansion of the calf muscles with anterior displacement of the neurovascular bundle. The suggested diagnosis was a soft tissue tumour such as a rhabdomyosarcoma. The chest X-ray was normal. The laboratory tests (full blood count, urea and electrolytes, liver-function tests and inflammatory markers) were essentially normal. The erythrocyte sedimentation rate was 24 mm/hr. The Mantoux skin test was normal. The bone scan showed no other areas of uptake apart from the left calf region.

Surgical exploration was performed under tourniquet control. The mass was well delineated arising proximally deep to the medial gastrocnemius, from the soleus muscle belly and extending distally to the tendinous junction between soleus and gastrocnemius. The muscles were stretched over the mass and the neurovascular bundle was stretched and adherent anterior to the mass, but dissected off easily. The mass was oval, well-defined, and rubbery in consistency with a pseudocapsule (Figure 4).

Figure 1. A 3-year-old child with diffuse swelling of the left calf

Figure 2. Anteroposterior and lateral radiographs show widespread speckled calcific density and opacities in the calf. There is no bone involvement.

Figures 3 (a) and (b). Coronal and axial MRI images with intravenous contrast show an oval soft tissue mass in the posterior compartment muscles of the leg with a heterogeneous signal intensity and demonstrating intense enhancement. There are areas of T2 and T1 hypointensity, which do not demonstrate enhancement post contrast in keeping with areas of ossification. There is expansion of the muscles with anterior displacement of the neurovascular bundle.
There was no increase in vascularity in the region and bleeding was minimal. The mass measured 9 cm × 5 cm. Macroscopically the specimen was firm, solid and had to be sectioned with a band saw and showed calcific flecks and multifocal areas of ossification. It was grey-white in colour, with a firm solid consistency. A gritty sensation was noted on sectioning.

Microscopic staining with haematoxylin and eosin (H&E) demonstrated a spindle cell tumour with large areas of hyalinisation, calcification and ossification. Necrosis was not seen. The tumour pattern was one of interlacing fascicles. The individual tumour cells contained clear and eosinophilic cytoplasm, elongated, blunt-ended nuclei with minimal size variation, and variable cytoplasmic vacuolation was evident. The mitotic count was <1 per 50 high power fields (HPF). Special stains confirmed a smooth muscle immunophenotype confirmed by alpha smooth muscle actin and desmin immunopositivity. The immunohistochemical markers for skeletal muscle, neural, epithelial, melanocytic, chondroid and endothelial lineage were negative. Necrosis, cellular pleomorphism, haemorrhage and vascular or perineural invasion was not seen. Entrapped aponeurotic and atrophic skeletal muscle tissue was seen. There was no atypia or features of malignancy. Epstein Barr virus encoded RNA testing was negative. The histopathological features suggested a myoid tumour compatible with a deep leiomyoma. The excision margins were free of tumour. The post-operative course was uneventful. The radiographs at follow-up were normal (Figure 5).

Discussion

Leiomyoma of deep soft tissue is exceptionally rare compared to its malignant counterpart. Leiomyoma is a benign, usually solitary tumour of smooth muscle origin commonly involving the organs of the body composed of smooth muscle such as the uterus, urogenital tract and gastrointestinal tract. When found in soft tissue, it usually involves the dermis and subcutaneous layer. Very rarely affects the deep soft tissue. The deep leiomyomas have been found most frequently within skeletal muscle originating from vessels with unstriated muscle. They are more common in adults and rarely described in children. The tumour usually shows calcification in 58% of cases. The largest tumour described in children measured 7.5 cm.

The radiological appearance can resemble various familiar conditions seen in orthopaedics and the diagnosis can be challenging.

Leiomyoma involving skeletal muscles of the extremities in children has been reported in nine children only, viz: thigh muscles = 3, arm = 1, calf = 2, deltoid = 1, pectoralis major = 1 and subscapularis = 1. Calcification occurred in six cases. Calcification with ossification was only seen once. A history of minor trauma was recorded in three patients.

Histologically, these lesions are composed of well-differentiated, smooth muscle cells with abundant eosinophilic cytoplasm similar to vascular smooth muscle. The existence of leiomyomas of deep soft tissue has been questioned. The tumours should be diagnosed using the most stringent criteria. They are highly differentiated smooth muscle tumours. Somatic leiomyomas should show no necrosis, at most mild atypia, virtually no mitotic activity with <1 mitosis per 50 HPF. Spindle cells are arranged in a parallel fashion with eosinophilic cytoplasm. The cells appear to originate from the outer layers of vascular channels. Tumour cells are usually positive for vimentin, desmin and smooth muscle actin.

Calcification is usually macroscopic and visible as opacities on radiographs. Microscopic calcification is also reported. The precise nature and pathogenesis of calcification in deep leiomyomas is uncertain. Classically, they have been considered to be dystrophic changes following tumour haemorrhage, necrosis or liquefaction. More recently, alkaline phosphatase, similar to uterine myoma, has been detected in the tumour, suggesting that the enzyme may play a role in the pathogenesis of calcification. Calcification may be scattered foci. In one report, the calcifications consisted of laminated, concentric spherules. In another, the calcifications appeared ‘mulberry’ in type, in the subscapularis muscle and the calf.
Calcific deposits in soft tissue can cause diagnostic challenges in orthopaedics. Among the conditions which produce localised soft tissue calcification are calcified haematomas, myositis ossificans, tumoral calcinosis, tuberculous lymphadenitis, pilomatrixomas, calcifying fibrous pseudotumour, synovial sarcoma, mesenchymal chondrosarcoma and extraskeletal osteosarcoma. These have similar radiological findings to leiomyomas in deep soft tissue.1,4

The calcified haematoma is composed of granulation and fibrous tissue, siderophages, inflammatory cells and coarse calcification. The granulation tissue reflects the reactive nature of the process. Myositis ossificans also occurs in muscle. Microscopically, it is well circumscribed and has a zonal pattern that is reflective of the degree of cellular maturation. The outermost zone is composed of mature, including lamellar, bone; and the innermost zone is comprised of a centrally fibroblastic tissue. The intermediate zone contains ill-defined trabeculae of admixed fibroblasts, calcification, osteoblasts, osteoid and ectatic vascular channels. The ossific appearance of the radio opaque mass and its conformity to muscle planes make the distinction from the popcorn-like calcification of leiomyomas. The morphology of tumoral calcinosis depends on the stage of the disease. In the early proliferative phase, foamy histiocytes, fibroblast-like cells and siderophages are seen. In the active stage, macrophages, giant cells, osteoclast-like giant cells characterised by fibrosis and calcification are seen. A smooth, muscle-rich component is not a feature of any stage. Fluid levels may be seen in the tumour. Calcified lymph nodes of tuberculosis demonstrate variable fibrocalcific alterations with stippled and coarse calcification. A variable palisade of histiocytes, granulomas and hyalination are seen.

Pilomatrixomas are tumours of hair follicle origin that are composed of sheets of basaloïd, epithelial cells and anucleate ghost cells. A spindle-cell component is not a feature of this tumour in the cellular or late calcified and ossified stage. These tumours are benign calcifying epitheliomas which occur in the skin and the subcutaneous tissue. Radiographically, they are well-delineated soft tissue tumours which contain fine sand-like calcification.11

Parasitic cysts due to hydatidosis and cysticercosis may calcify. While granulomas, histiocytes and eosinophils may be variably present around the calcified cyst wall, there is no evidence of an associated smooth muscle proliferation.

Scleroderma, calcification due to calcium gluconate extravasation, haemangiopericytoma, fibrosarcoma and ossifying lipofibroma can also produce localised soft tissue calcification.

Calcifying fibrous pseudotumours are hypocellular, hyalinated, fibrosclerotic tumours with psammomatous calcification. There is a variable inflammatory background with or without foamy histiocytes. It contrasts with the cellular nature of leiomyoma.

Monophasic synovial sarcoma shares the spindle-cell morphology of leiomyoma. There may be fine coarse calcification and ossification. However, the cells lack the eosinophilic cytoplasm that is characteristic of smooth muscle cells. In addition, synovial sarcoma expresses variable epithelial membrane antigen and cytokeratin immunopositivity.

Extraskelatal osteosarcoma contains pleomorphic osteoblasts in osteoid. There is hypercellularity, increased mitotic activity and necrosis. Leiomyoma lacks pleomorphism, necrosis, high mitotic activity and malignant osteoblasts.

Mesenchymal chondrosarcoma is a biphasic tumour with sheets of undifferentiated spindle, round and oval cells, a haemangiopericytic pattern and lobules of hyaline cartilage. Focal calcification and ossification of the cartilaginous component may be present. Leiomyoma lacks these characteristics.

Epstein Barr virus (EBV)-associated smooth muscle tumours are a recently emerging entity that occur in the setting of immunocompromised patients at a greater frequency than the general population.2 Smooth muscle tumours have now been reported in patients with various types of organ transplants in patients with Aids and in those with congenital immunodeficiency.2 The common denominator in all cases appears to be EBV virus infection in the setting of immunocompromise. EBV can be detected by in situ hybridisation in virtually all tumours studied.2 In these patients the tumours are mainly visceral. In a five-year clinico-pathological review, Ramdial et al described six leiomyomas in patients with Aids associated with the EBV to increase the awareness of this association. Two of these children had masses in the dermis of the lower limbs. They also had associated tuberculosis of the chest. The intramural EBV status of all tumours was determined by EBV-encoded RNA in situ hybridisation.16,17

Conclusion

The outcome of treatment of leiomyoma of deep soft tissue is usually good. In previously reported cases of leiomyoma, total excision of the tumour provided complete cure without recurrence. Leiomyomas very rarely undergo malignant transformation.1,4,5 The location of the tumour reported in this study is uncommon. The progressive calcification with ossification is unique and important in differential diagnosis. A well-defined circumscription with a pseudocapsule is a characteristic feature.

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References

KEY WORDS REQUIRED IN ARTICLES

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