Glomus tumor of digital artery of thumb – Case Report

Glomangioma da artéria digital do polegar – Relato de um caso

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Abstract
We report on a case of glomus tumor in the branch of the radial artery of the right thumb. The patient had a 4-year clinical history. The patient reported the development of a 2.0-cm tumor in the interdigital region between the first and second fingers of the right hand on the short flexor muscle of thumb. The patient also complained of severe and progressive pain. Tests using bidirectional Doppler and echo-color-Doppler revealed a presumptive diagnosis of arteriovenous malformation based on the turbulence of the flow and absence of stenosis. The tumor was removed by open surgery and sent for histopathological examination, which showed a diagnosis of glomangioma. The present report describes a rare arterial disease causing extreme discomfort to the patient, which may be treated with surgical resection without sequelae.

Keywords: glomus tumor; glomangioma; radial artery.

Resumo
Os autores relatam um caso de tumor glômico em ramo da artéria radial que irriga o polegar direito, com história clínica de 4 meses. O doente referia o aparecimento de tumoração de aproximadamente 2,0 centímetros na região interdigital, entre o primeiro e o segundo quirodactilos da mão direita, sobre o músculo flexor curto do polegar, extremamente dolorosa e com dor em progressão. Os exames com aparelho de Doppler bidirecional e o eco-color-Doppler apresentaram, como diagnóstico presuntivo, malformação arteriovenosa, pelo turbilhamento do fluxo e ausência de estenoses. A tumoração foi retirada por cirurgia aberta e encaminhada para exame histopatológico, com diagnóstico de glomangioma. Este relato descreve uma doença arterial pouco frequente, que causa extremo desconforto ao seu portador, mas que é solucionada pela exerese cirúrgica, sem sequelas.

Palavras-chave: tumor glômico; glomangioma; artéria radial.

Introduction
The glomus tumor is a benign familial disorder. It originates from the glomus body, is often smaller than 3 mm and is formed by nerve, muscle and arterial components. This rare tumor accounts for less than 5% of the hand tumors and is more frequent in young women in their 20 to 40 years1. It is often found in subungual regions, and its clinical presentation includes local hemorrhage. Its main characteristic is intense local pain. The diagnosis is made according to clinical symptoms, and only one third of the cases show radiological abnormalities, such as bone erosion2-7. The differential diagnosis should include other hand tumors, such as neurilemmoma, hemangioma, osteoma and schwannoma. Surgical removal results in cure and immediate pain relief.

Case report
A 58-year-old man presented with a tumor with a 4-month history, measuring about 2 cm, extremely painful, located in the dorsal region of the right hand, between the first and second fingers over the short flexor muscle of the thumb. He denied any local trauma, exhaustive physical activity or any other probable cause of tumor formation. His pain worsened gradually, and he had an enlarged
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Operative technique and results

The patient underwent regional anesthesia of the right upper extremity using 1% lidocaine, with no vasoconstrictive agents. A skin incision was made, tissue layers were separated, and the vessels were carefully dissected. A large number of collateral veins were seen, and some had thrombosis. The tumor was encapsulated in the artery, which was fully removed, and the pseudocapsule was preserved and sent to the laboratory for histopathology examination.

Immediately after operation (24 hours), pain disappeared. At follow-up, the patient was asymptomatic and had returned to work.

Histopathology

Tumor made up of polygonal cells with small and regular nuclei, some in solid groups, but others in regularly organized cell threads. Few mitotic figures were found. Dilated and congested blood capillaries proliferated in the middle of the tumor cells, containing recent thrombi and covered by a single layer of endothelial cells (Figure 2).

Discussion

A neuromyoarterial glomus and its tumor were first described by Masson in 1924, and in 1972 Martorrel classified it as a glomangioma, or glomus tumor. It is formed by nervous fibers, muscle cells and vascular components and, for that reason, it was called glomangioma.

It is a rare tumor and accounts for less than 2% of all tumors that affect soft tissues. Its appearance and clinical history almost always involve some type of trauma. Pain...
is reported by all patients. In the beginning, pain is mild, but later becomes intense and unbearable. There is some regularity in time between pain episodes, but, as the disease progresses, pain frequency increases and the simple touch of clothing is often enough to trigger a crisis. Pain usually occurs at night, which makes it impossible for the patient to sleep. One of the hypotheses to explain pain intensity is based on tumor expansion, that is, as it is contained in a pseudocapsule, its growth is restricted, which leads to the necrosis of the central cells and their replacement by connective tissue. Another hypothesis to justify pain is tumor site, as it arises in the myoneural junction.

The region most frequently affected by an extremity glomangioma is the dermis, the subcutaneous cell tissue, particularly in the nail bed, because there is little resistance to its development there, and not in small-caliber arteries, as Kamarashev et al. used to believe. These tumors may also be found in other sites, such as the stomach, knees, shoulder and mediastinum, as well as in the middle ear, where it may lead to serious balance disorders and hearing loss. They are never larger than 5 mm, regardless of the region where they arise or its progression time. When it is subungual and does not have space to grow, it loses its oval shape and may cause bone erosion, which may give the false impression of a malignant infiltrating tumor.

Imaging studies, such as arteriography and echo-color Doppler ultrasound, are useful to evaluate glomangiomas of the extremities, particularly to make the differential diagnosis with other tumors, but images are not conclusive. Its diagnosis, in general, is made early due to pain, but studies in the literature describe cases that had a history of 40 years. Some authors suggest that cell atypia is a result of the accumulation of heterochromatin associated with DNA inactivation. Moreover, the increased capillarity, the local architecture in the arrangement of tumor cells with uniform sizes and forms, and the uniformity of base cells suggest the diagnosis of glomus tumor. From the moment the diagnosis of a glomangioma is made, no other case of such tumor goes unnoticed by the examiner because of its particular characteristics.

Treatment is surgical using a conventional or laser technique, and the latter is applied in special to the cases of multiple glomangiomas. However, there may be tumor cell implantation the capsule ruptures, and a new tumor may be implanted, at a mean recurrence rate of 13.3%. Tumor resection provides immediate symptom relief and complete pain resolution. In this case, we chose an open surgery because total tumor removal was important to avoid the rupture of the capsule that contained it, the implantation of glomus cells and tumor recurrence, in agreement with treatment data in the literature.

References


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