CASE REPORT

Fine Needle Aspiration of Glomangioma Knee - A case report

Syed Besina Yasin, M.D., Sumyra Khurshid Qadri, M.D., Nassima Chanda, M.D., Rumana Hamid Makhdoomi, M.D.
Department of Pathology, Sher-i- Kashmir Institute of Medical Sciences (SKIMS), Srinagar (J&K), India

ABSTRACT

Glomus tumor, a benign neoplastic proliferation of modified smooth muscle cells characteristically occurs in a digital subungual location and presents with localised pain. Extra-digital locations that have been described so far include bone, tongue, stomach, rectum, lung, mediastinum, sacrum, coccyx and head and neck areas. Though the histology of glomus tumor is very characteristic, cytological features are poorly defined. Less than ten cases describing cytologic features have been reported so far. We present a case of glomus tumor of the knee joint detected on fine needle aspiration and confirmed later on histopathology, in a 55-year-old male who presented with history of painful nodule on the anterior aspect of the left knee joint for one year. JMS 2011;14(2):71-73

Keywords: Glomangioma, fine needle aspiration cytology

Case report

A 55-year-old male presented with history of painful nodule on the anterior aspect of the left knee joint for past one year. On physical examination, there was a small 0.5 cm subcutaneous nodule. It was reddish-blue in colour, mobile, firm and was intensely painful on touch. Routine investigations were within normal limits. A clinical diagnosis of neurofibroma or glomangioma was made. Fine needle aspiration cytology (FNAC) was performed using a 20 ml syringe with 22 gauge needle. The patient experienced excruciating pain during the procedure. A blood-mixed material was obtained. Smears were stained with Papanicolaou and May-Grunwald Giemsa stains. On microscopic examination, the aspirate was heavily admixed with blood. There were cohesive clumps of uniform round to oval cells along with wisps of magenta coloured intercellular material. The nuclei were round to oval, with firmly distributed chromatin. Intranuclear inclusions, nucleoli and mitosis were not seen. Cytoplasm was poorly defined (Figure 1). A few groups of spindle shaped cells were seen near the clusters of tumour cells. Numerous RBC’s were seen in the background. A diagnosis of glomangioma was offered on FNAC based on the clinical features and the cytologic features.
The lesion was excised and gross examination revealed a small nodule 0.5 x 0.5 cm in size. The cut surface was greyish white. Microscopy revealed a well circumscribed tumor consisting of uniform round to oval cells in nests and clumps around thin walled capillary sized blood vessels which were lined by endothelial cells. Cells showed a uniform central rounded nucleus with mild to moderate eosinophilic cytoplasm (Figures 2 & 3). PAS and reticulin stain demonstrated a dense reticulin network investing the single tumor cells. Based on all the above features, a diagnosis of glomus tumor was given. Postoperatively, the patient was completely relieved of pain.

**Discussion**

Glomus tumors are uncommon tumors with an estimated incidence of 1.6% among the 500 consecutive soft tissue tumors reported from the Mayo Clinic. They are composed of cells that resemble the modified smooth muscle cells of a glomus body. Normal glomus cells are located in the subcutaneous tissue which is responsible for the regulation of temperature and blood pressure. Glomus tumors may be single or multiple. Usually, single tumors are idiopathic and multiple ones are inherited as an autosomal dominant trait. The tumor is about equally common in both genders, although there is a striking female predominance among patients with subungual lesions. Most glomus tumors are diagnosed during adult life. The classic triad of aching pain, exquisite tenderness and temperature sensitivity provide the diagnosis. The most important site is the subungual region but other common sites include palm, wrist, forearm and foot. Rare sites include stomach, mediastinum, lung, trachea, vagina, etc.

Our patient was a 55 year male with a solitary 0.5 cm diameter subcutaneous nodule on the anterior aspect of the left knee which is a less common site. Patient presented with aching pain and extreme tenderness. A clinical diagnosis of glomus tumor / neural tumor was offered. There was no clue from the radiological investigations to confirm the diagnosis. FNAC was done. A diagnosis of glomus tumour was suggested on the basis of typical features on FNAC and later confirmed on HPE of the excised tumour mass.

Less than ten cases of this tumor have been reported on fine needle aspiration. Cytological features of this tumor were first described by Holck and Bredeson in 1996. They observed that the aspirated smears were cellular, composed of tight clusters and more loosely packed sheets of medium sized polygonal cells as well as frequent stripped nuclei. At places, the cell groups form a 3-dimensional pattern. The nuclei were round to oval with finely distributed chromatin. Intranuclear inclusions and nucleoli were inconspicuous as were mitosis. The cytoplasm was poorly defined. Rows of spindle cells, probably endothelial cells, focally crisscrossed
the cell groups. At places delicate feathery, curved threads of reddish material corresponded to myxoid substance and basement membrane material was seen.²

Vidyavathi et al, reported a glomus tumor on fine needle aspiration of a 28-year-old female with a 2 x 1 cm mobile swelling in the forearm.¹ Endoscopic ultrasound guided fine needle aspiration of gastric submucosal tumors has revealed one glomus tumor in a study conducted by Sawaki et al, on 304 lesions. Vinette and Yazdi have also reported a glomus tumor of stomach on fine needle aspiration in a 72-year-old asymptomatic female.³ M Perez-Guillermo et al, in a study of 12 cutaneous vascular tumors by means of fine needle aspiration, reported one glomus tumor⁴. The differential diagnosis includes cutaneous adnexal tumors like eccrine spiradenoma, carcinoid tumor or a hemangiopericytoma.⁵ Due to location, adnexal tumor was the differential diagnosis in our case. Cytologic features of glomus tumor and eccrine spiradenoma which form close differential diagnosis have been rarely reported. Smears of eccrine spiradenoma show presence of bland uniform cells in cohesive clusters and cribriform sheets with rosette-like structures surrounding amorphous material. Three types of cells are also seen-large epithelial cells, myoepithelial cells and smaller lymphocytes.¹ All these features were not seen in our case.

The presence of excruciating pain on aspiration and cohesive clusters of small round cells with presence of basement membrane like magenta-coloured material along with groups of spindle shaped endothelial cells lead to the diagnosis of glomus tumor.

The histologic features of glomus tumor have been well described in literature. On gross examination, they are blue or purple and less than one centimetre in size. Microscopically, the glomus tumors are relatively well circumscribed capsulated lesions and show numerous small vascular lumina surrounded by clusters of monotonous polygonal glomus cells set in a hyalinised or myxoid stroma. The glomus cells are characterised by a faintly eosinophilic cytoplasm and large, punched out pale nuclei with delicate chromatin and inconspicuous nucleoli.⁶ On Immunohistochemistry, vimentin and SMA can be identified in nearly all glomus tumors. Desmin is highly variable.¹

In conclusion, it is emphasised that in addition to neural tumors and adnexal lesions like eccrine spiradenoma, glomus tumors should also be considered in the differential diagnosis of all soft tissue tumors associated with pain and tenderness. Fine needle aspiration is a good tool at arriving the correct diagnosis in such cases.

References