Case Report

The mysterious chronic finger pain- glomus tumour: a case report

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ABSTRACT

Glomus tumours are very rare, small, painful benign hamartomas, arising from the arterial end of the glomus body. Glomus tumors account for 1-2% of the soft tissue tumors in the hand. The exact etiopathogenesis is unknown. It is incapacitating to the patients because of chronicity of symptoms and lack of proper investigative tools to diagnose the tumor at an early stage. The clinical diagnosis was made on the basis of medical history and MRI findings. We report a case of glomus tumor in a 35-year-old female, situated in the pulp of distal phalanx of right middle finger, which was resected completely. Excision of lesion imparted complete resolution of symptoms. The histopathological report was consistence with glomus tumor.

Keywords: Finger, Glomus tumour, MRI hand, Pain

INTRODUCTION

The normal glomus body is an arteriovenous shunt involved in thermoregulation, mostly concentrated in the digits. Glomus tumors are rare benign hamartomas composed of endothelium lined vascular spaces (Sucquet-Hoyer Canal) surrounded by glomus cells.1,2 Arising from neuromyoarterial apparatus of glomus body,7 Hoyer first described them in 1877, while the first complete clinical description was given by Masson.4

 Majority of glomus tumors are small benign neoplasm that occur in the dermis or sub cutis of extremities.5 Glomus tumors can be solitary or multiple. Some cases of large multiple visceral lesions are reported to be malignant.4 The précised etiopathogenesis is unknown.8

Average age at presentation is from 30-50 years of age. Glomus tumors are usually under 1cm in size, present as a faint bluish purple papule and associated with classic triad of symptoms: local sensitivity, pain with cold exposure and severe pain on minor trauma.2

CASE REPORT

A 35-year-old Hindu female came to surgical clinic with history of pain at the tip of right middle finger for the last 2-years. She denied history of any preceding trauma. She had worsening of symptoms at coughing or sneezing, but there is no history of root pain. She had localised pain to a point on the right middle finger near its tip over distal phalanx. There is no history of any skin discoloration. She gave history of worsening of pain on cold exposure, but no history of a typical of Reynaud’s phenomenon. She gave history of worsening of pain on cold exposure, but no history of a typical of Reynaud’s phenomenon. On examination there was sharp localised tenderness over the right middle finger near tip. There was no local rise of temperature or any discoloration. She was afebrile and no any regional lymphadenopathy was detected. Routine blood examination and X-Rays right hand were within normal limits. HRMRI proved the diagnosis of Glomus tumour (Figure 1). Complete excision was done and patient became fully asymptomatic following the surgery and there were no postoperative complications. No recurrence of symptoms was found at one year follow up. Resected specimen sent
for biopsy (Figure 2) and report was consistent with the diagnosis of Glomus tumour (Figure 3).

Figure 1: MRI mass with intense high signal on T2 image.

Figure 2: The mass removed en block.

Figure 3: Neoplastic cells with monomorphic round or oval nuclei and abundant pink cytoplasm. Tumour cells invading vessel wall.

DISCUSSION

Glomus tumour is a slow growing benign lesion with intense pain. Lesions are characterised by a faint bluish purple papule, paroxysmal pain, pin head tenderness and cold sensitivity. The lesions are usually located on extremities, especially in subungual area, followed by lateral aspect of digits, pulp and palm, but may occur on any other parts of body. Glomus tumour is a slow growing benign lesion with intense pain. Lesions are characterised by a faint bluish purple papule, paroxysmal pain, pin head tenderness and cold sensitivity. The lesions are usually located on extremities, especially in subungual area, followed by lateral aspect of digits, pulp and palm, but may occur on any other parts of body. Multiple glomus tumors are common in male called glomangiomas and have autosomal dominance pattern with incomplete penetrance. Rarely glomus tumors may present with clinical features like large size, visceral origin (GUT and GIT), multi-centricity and infiltrative growth pattern. The delay in diagnosis causes immense morbidity to patients. Proper history, high index of suspicion and careful clinical examination are prerequisite to clinch the diagnosis. Routine laboratory investigations have no role in diagnosis of glomus tumor. High resolution Magnetic Resonance imaging (HRMRI) is the investigation of choice and assesses tumour characteristics more accurately (Figure 1). High resolution CT scan and X-Rays were not helpful in our case. The treatment of choice is complete excision, which is crucial in prevention of recurrence and reduction of symptoms, with low rate of recurrence. A bloodless surgical field is essential to allow meticulous removal of tumor. We excised the tumor completely and there was no recurrence of pain on follow up for 1 year. Complete excision provides best chance of cure with best functional outcome. The diagnosis was confirmed by histology (Figure 3). The abatement of symptoms following surgery is quite pleasurable both for the patient and the surgeon.

CONCLUSION

Accurate preoperative localisation of glomus tumour is vital for complete extirpation and to avoid recurrence. HRMRI is the investigation of choice. Complete excision ensures best chance of cure with best functional outcome. The alleviation of symptoms following surgery is quite gratifying both for the patients and the surgeon.

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