Calcified Vocal Cord Nodule – A Unique Case Report

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ABSTRACT
Vocal cord nodules are benign neoplastic lesions which occur due to submucosal oedema and haemorrhage, leading to fibrosis and hyalinization. Calcification in vocal cord nodules has not been reported so far in literature. It is thought to be a laryngeal counterpart of idiopathic calcinosis cutis. Here, we are reporting a case of a 38-year-old male patient who presented with a change in voice, which had a duration of one month. Laryngoscopic examination revealed a globular, yellowish white, sessile mass which arose from anterior commissure region of right vocal cord. Micro laryngeal excision was done. Histopathological examination was suggestive of a calcified nodule in vocal cord.

On haematoxylin eosin staining, the microscopic section showed stratified, squamous epithelium overlying stroma, which contained sub epidermal collection of calcium deposits which were seen in dark blue stains. Post-operatively, serum free calcium, parathormone and calcitonin levels were studied and they were found to be normal. Patient was followed up for a period of 6 months. He was symptom free and laryngoscopic examination revealed normal vocal cords.

DISCUSSION
Vocal cord nodules are more commonly found in young women and in people who routinely stress their vocal cords (singers, auctioneers, hecklers). Vocal cord nodules are initially oedematous, with a myxoid stroma, but they will become fibrotic with time. Vascular proliferation may be seen with dilated vessels, granulation tissue and haemorrhage [1]. In our case, patient was a carpenter by occupation and there was no history of vocal abuse.

The subepidermal calcified nodule, also known as cutaneous calculi, is a form of idiopathic calcinosis that affects children [2]. It occurs in facial skin usually and has no relationship with connective tissue of vocal cord.
tissue disease or with any abnormality in calcium and phosphorous metabolisms. The oral cavity is very rarely affected by calcinosis cutis of any type. Two cases of calcified nodules were reported, one in gingiva of a 1-year-old girl and the other in the tongue of 5-year-old boy [2]. In our case, the patient was 38-year-old at the time of presentation and the site of involvement was vocal cord.

The sub epidermal calcified nodule presents as an asymptomatic, solitary, yellow-white or erythematous filiform tumour. The most common location is on the head and neck regions of male children [3] and a asymptomatic sub epidermal calcified nodule in a child's sole has also been reported [4].

Calcified nodules have been reported in thyroid, lung, abdomen and brain. In our case, patient presented with a calcified nodule in vocal cord, but no reference regarding calcification of vocal cord nodule has been reported in literature so far. Calcification of soft tissue can be caused by vitamin K2 deficiency or by poor calcium absorption, due to a high calcium or vitamin D ratio. This can occur with or without a mineral imbalance. Abnormalities in calcium-phosphate balance develop early in the course of chronic kidney disease [5].

A variety of factors can cause this condition; the most common cause is dystrophic calcification which occurs in soft tissue as a response to injury, inflammation, infection, tumours, diseases of connective tissue, hypocalcaemia, hyperphosphatemia. The causes and histiogenesis of these deposits have not been fully established, but the most favoured theory is that calcium is deposited in a pre-existing lesion. The treatment of choice is surgical removal, with a histopathologic examination [2].

In idiopathic cutaneous calcinosis, histopathologic examination reveals a focal papillary dermal collection of dark blue –staining, large amorphous deposits and or small calcified globules which are surrounded by a lymphohistiocytic infiltrate [3]. The causes of histiogenesis of these deposits have not been fully established, but most favoured theory is that calcium is deposited on a pre-existing lesion. In our case, on histopathological examination, the section showed stratified, squamous epithelium overlying stroma, which contained sub-epidermal collection of calcium deposits which were seen in dark blue stains.

Due to the rarity of its site, we consider this case to be unique and special attention should be given to calcification of vocal cord nodules among other benign neoplastic lesions of vocal cord.

**CONCLUSION**

This case has been reported because there is no substantial literature which has revealed calcification which occurred in vocal cord. Therefore, our case is unique in showing an unusual site of involvement.

A calcified vocal cord nodule has to be considered as a differential diagnosis for benign neoplastic lesions of vocal cord.

**REFERENCES**


