Case Report

Post Traumatic calcinosis cutis of eyelid

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Introduction

Initially described by Virchow in 1855, calcinosis cutis is an uncommon disorder characterized by an abnormal deposition of insoluble amorphous calcium salts under the epidermal layer of skin in various parts of body (Reiter N et al, 2011). This syndrome is divided into four types: dystrophic calcification, metastatic calcification, idiopathic calcification and iatrogenic calcification. The dystrophic type presents in damaged and devitalized tissue. Metastatic calcification presents in systemic diseases. Iatrogenic type is associated in many cases with intake of calcium containing medications. Idiopathic type occurs in absence of any detectable cause or tissue injury (Reiter N et al, 2011). Here we describe a rare case of post-traumatic calcinosis cutis of the eyelid.

Case

A 12 year old male child presented in the eye department of our hospital with complaint of a grayish mass on the lateral aspect of left upper eyelid for one year (Figure 1). A significant history of trauma one and a half year back with a pencil tip while playing with his friend was obtained. The trauma got neglected due to minor pain and no major treatment was undertaken at that time. There was no significant systemic or family history. On physical examination, child was moderately built. No other masses were found in any other part of his body. There was no lymphadenopathy. On ocular examination, best corrected visual acuity was 6/6 for both eyes (Snellens chart). Anterior segment examination in right eye was within normal limits. In left eye, a grayish pigmentation was noted just below the eyebrow of around 4mm size which was not associated with redness or vascularisation. On palpation, a hard consistency was felt. The mass was non tender and non pulsatile. Rest of the examination of anterior and posterior segments was within normal limits. On the basis of history and examination, a provisional diagnosis of post traumatic left upper eyelid retained foreign body (most probably lead from pencil tip) was made. All routine haematological investigations including serum calcium and phosphorus were within normal limits. Excision of the mass was planned under local anaesthesia.

A superficial incision was made over the mass, but since it was deeper in the dermis so incision had to be extended (Figure 2). The nodule was removed en-masse. It was 7mm in size (Figure 3). The mass was sent for histopathological examination. Histopathological findings showed that the specimen was lined by stratified squamous epithelium. The underlying dermis showed a large area of calcification. The area was surrounded by fibrous tissue and few chronic inflammatory cells. All these features were suggestive of calcinosis cutis (Figure 4 a and b). There was no trace of any retained lead or pencil tip. It has been more than 6 months and no recurrence has been noted.
Figure 1: Per-op photo of the child. Arrow directed towards the mass.

Figure 2: Intraoperative photograph showing that the mass was deep into the dermis.

Figure 3: Mass as taken out en-masse (7 mm in size)

Figure 4 a and b: Histological findings show that the specimen is lined by stratified squamous epithelium (A). The underlying dermis shows a large area of calcification. The area is surrounded by fibrous tissue and few chronic inflammatory cells (B). Calcium deposits in the dermis can be seen (C). All these features are suggestive of calcinosis cutis.

Discussion
To our knowledge and after thorough literature search, this is the first case report of a patient with post traumatic calcinosis cutis of the eyelid. Though there have been reports of idiopathic calcinosis cutis published previously in the literature. Samaka et al, 2015 reported a case of idiopathic calcinosis cutis in upper eyelid. Malik et al, 2011 reported a case with idiopathic bilateral subepidermal calcific
nodules in both eyelids. In idiopathic form it may mimic as molluscus (Nguyen J, 2008) or solitary verruca nodule (Nico MM, 2001). Idiopathic calcinosis cutis of the tarsus of upper eyelid in a patient with Rheumatoid arthritis with normal serum calcium levels has been reported (Ikhyun Jun, 2011). It has been seen to have an association with disorders like dermatomyositis, systemic lupus erythematosus or systemic sclerosis (Boulman N, 2005). Eyelid, palpebral and bulbar calcifications also have been noted in patients with inflammation, systemic hyperparathyroidism and hyperphosphatemia (Ferry AP et al, 1990 and Ghanchi, F et al, 1996). In our case serum calcium and phosphate levels were within normal limits and no other cause could be found out except a clearly defined history of trauma at that exact point. Investigations should be done to exclude any underlying pathology especially in idiopathic cases.

Histological examination of the lesion reveals calcium deposits in the dermis in biopsy, which may or may not be surrounded by foreign body giant cell reaction. No standard treatment has been recommended for the removal or reduction of the lesion of calcinosis cutis (Boulman N et al, 2005 and Dutz J et al, 2005). Colchicine, warfarin, bisphosphonates, probenecid and diltiazem have been used with varying degree of success (Dutz J et al, 2005). Surgical excision is the treatment of choice (Ikhyun Jun et al, 2011).

Conclusion
This is the first case report of post-traumatic calcinosis cutis affecting the eyelid which is a rare site for involvement.

References


