PAROTITIS IN LIPOID PROTEINOSIS

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Abstract

Lipoid proteinosis is a rare autosomal recessive multisystem disorder caused by mutations in the extracellular matrix protein 1 gene. It is characterized by infiltration of amorphous hyaline in the dermis and submucosal connective tissue. We report the case of a 36-year-old woman with history of hoarseness since infancy and multiple episodes of right parotitis. Dermatological examination found bead-like papules on the margins of the eyelids, thickened skin lesions on the elbows and knees, enlargement of the tongue with dental impressions and lingual frenulum infiltration. Skin biopsy showed histopathological findings of lipoid proteinosis. The patient was treated by acitretin for 6 months with no improvement of the skin lesions. Three months later, patient presented a left parotitis with a cutaneous fistula through which abscess was discharged. The objective of the present report is to describe this rare entity and report that lipoid proteinosis may be associated with recurrent parotitis caused by infiltration and stenosis of Stensen’s duct.

Introduction:--
Lipoid proteinosis (LP) is a rare genodermatosis characterized by infiltration of amorphous hyaline in the dermis and submucosal connective tissue (Bhattacharjee et al., 2018). It was first described by the dermatologist Urbach and the otolaryngologist Wiethe in 1929 (McGrath, 2015).

Case report:--
A 36-year-old female born of a consanguineous marriage with history of hoarseness since infancy and multiple episodes of right parotitis occurring within a six years period.

Cutaneous examination revealed multiple beaded papules lining the free edge of eyelids (Fig 1.a), verrucous papules over the dorsum of hands, hyperpigmented scars over elbows and knees (Fig 1.b) and a mild frontal alopecia (Fig 1.c).

Lips were infiltrated and commissures presented small waxy papules. Examination of the oral cavity showed some white papules on the inner surfaces of the lips with a hypertrophy of Stensen’s duct orifice (Fig 2.a). The tongue was hypertrophied and showed dental impressions (Fig 2.b). The lingual frenulum was thickened and seemed to impair tongue mobility (Fig 2.c). The clinical findings along with the histopathological examination of skin supported the diagnosis of LP. No neuropsychiatric abnormalities were reported. MRI of the head has not revealed calcifications. Laryngoscopic examination and ophthalmologic evaluation were within normal limits.

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Acitretin was started at 0.5 mg/kg/day. After 6 months of treatment, no improvement of the skin lesions was noted, and the treatment was discontinued.

Three months later, patient presented an increasingly painful swelling and reddening in the area of the left parotid gland. A cutaneous fistula was developed through which abscess was being discharged (fig 3.a).

Ultrasound of parotid glands revealed a swelling of the left gland with duct dilatation and heterogeneous parenchyma with a hypoechoic zone in deep lobe. No image of calculus was observed.

Antibiotic therapy was given for 10 days. Subsiding the swelling and pus discharge. It persists a scar tissue on the left cheek on anterosuperior region of the angle of mandible. A small pinpoint opening was seen just above the scar from which watery fluid discharged spontaneously.

A week later healing was complete. The skin orifice has disappeared leaving behind trivial scarring. There was no residual swelling, no pain and no recurrence of the fistula. The patient was referred to Ear Nose and Throat department (fig 3.b).

**Discussion:**

LP is an autosomal recessive disorder caused by mutations in the extracellular matrix protein 1 gene (ECM1) located on chromosome 1q21. ECM1 is expressed in several tissue. It is involved in extracellular matrix formation and its mutation is responsible for deposition of non-collagenous proteins and glycoproteins in the skin and mucosa (Pinna et al., 2000; Ghazawi et al., 2019).

The most consistent clinical feature is hoarseness of voice due to laryngeal infiltration, often from birth or infancy (Larkin et al., 2018). The classical cutaneous symptom is moniliform blepharosis; it is a pathognomonic sign characterized by presence of multiple papules that line up on the eyelid margins (Belliveau et al., 2015).

Infiltration and thickening of oral mucosa may lead to upper submandibular gland inflammation as well as recurrent episodes of parotitis (Al-Ekris and Al-Sadhan, 2012), caused by infiltration and stenosis of Stensen’s duct, and lead to xerostomia (Epple et al., 2018). A dry mouth and repeated trauma owing to macroglossia may lead to the development of lingual ulcer (Sabater-Abad et al., 2019).

There is no curative therapy available for LP. Various treatment alternatives including acitretin, D-penicillamine, laser treatment have been tried in literature with no uniformly success (Dertlioglu et al., 2014).

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**Figure 1:** (a) Moniliform blepharosis. (b) Elbow skin thickening with hyperpigmented scars. (c) Frontal alopecia.
Figure 2: (a) Hypertrophy of Stensen's duct orifice. (b) Enlargement of the tongue with dental impressions. (c) Lingual frenulum infiltration.

Figure 3: (a) Left parotitis with swelling in the preauricular region. (b) Scarring after disappearance of the cutaneous fistula.

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